Practical Physiological Chemistry

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Thirteenth Edition

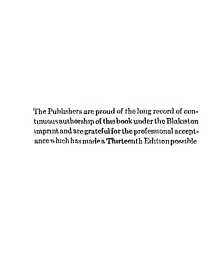
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Dedicated

To the memory of

Guiding Gennis of American Biochemistry who fifty vears ago inspired the First Edition of this work



Preface

It is with a mingled sense of pride and humility that the senior author presents this—the Thirteenth Edition of PRACTICAL PHYSIOLOGICAL CHEWISTRY To have been actively engaged in the authorship of successive editions of a textbook for fifty years is a privilege granted to few in the anaals of medical publication For this good fortune, I am deeply grateful

Perhaps even more than some of its predecessors, this Edition bas undergone extensive revision and expansion The proportion of the book devoted to textual material, as distinguished from experiments and procedures, has increased to the extent that it now accounts for about onehalf of the entire volume Certain ebapters, such as those dealing with Auclere Acids and Nucleoproteins (Chapter 7), Milk (Chapter 8), Muscular Tissue (Chapter 10), Enzymes and Their Action (Chapter 12), and Isotopes (Chapter 32), have been exhaustively rewritten To other chap ters major additions have been made, as for example Chapter 18 to which a section on Liver Function has been added, and Chapter 23 which now contains a section on Microclinical Chemistry

Try as its author will no textbook in such a rapidly developing area of research as biochemistry, can possibly be kept strictly up to the minute, it must nevitably run a losing race with the periodical literature. Nevertheless the authors of this volume and their collaborators have expended every effort in the direction of supplementing established knowledge with current facts, theories, and hypotheses, insofar as this is possible with full awareness that in some respects, obsolescence commences with the date of Publication (if not before) This may be of more concern to the research worker than to the student and for that reason the former should have recourse to original journals such as those eited profusely throughout this book

An endeavor has been made to keep pace with current advances in the many fields of biochemistry which are benefiting from the availability of advanced techniques and instrumentation Symbolic of these innovations, which give promise of epoch making advances, is the replacement of the time-honored Frontispiece of this book depicting absorption spectra of the blood pigments and their derivatives with a new color plate showing a two-dimensional chromatogram of a protein hydrolyzate For this illus trained the authors are greatly indehted to Dr A J P Martin of the Vational Institute for Medical Research, London and to the publishers of E. of Endeavour, The Imperial Chemical Industries Ltd, London

Experimental, analytical, and preparative methods employing these newer techniques and research tools are scattered through various chapters of the book Only a brief summary can be given here. The student and teacher of biochemistry will find of general interest the new material on electrophoresis the ultracentrifuge ion exchange resins, column and

paper chromatography; countercurrent distribution; the helical structure of proteins; the polypeptide sequence of insulin; the chemistry of the corticosteroids; the synthesis of oxytocin; the chemistry and metabolism of nucleoproteins, nucleic acids and derivatives; the role of muscle proteins and ATP, in muscle contraction; the hiochemistry of bone and teeth; the kinctucs of enzyme action, theories of blood clotting; the intermediary metabolism of earthohydrates, fats, and amino acids, including the role of ecenzyme A and "one-carbon fragments"; the cobalamins; thioctic acid, leucovorn, and related factors; isotopes and their use in biochemical research, new material on antibiotics; and much more.

The teacher will find many new experiments suitable for class use, some of which were developed specifically for this Edition, illustrating such topics as paper and column chromatographic separation of amino acids, purne nucleotides, etc.; countercurrent distribution; myosin and actin; glycogen storage and depletion; determination of blood volume with isotope-labeled red blood cells; blood coagulation and prothrombin time;

liver function tests; and many others

The chincian will find new and authoritative sections on: the biochemistry of liver disease; the biochemistry of the bones and teeth, and the use of fluoride in the prevention of dental decay; isotopes and their use in medicine; dextran; cholmesterase; adrenal cortical and pituitary hormones, and the nutritive value of milk. Chinical chemical procedures not found in the previous Editions include the more important liver-function tests (thymol turbidity, cephalm-cholesterol flocculation, hromsulfalcin, etc.); determination of blood protein-bound iodine, of blood cholmesterase, and of blood sodium and potassium by flame photometry; and others A unique and valuable section has also heen added on the use of micromethods in clinical chemistry, with detailed procedures' and descriptions of apparatus

Revisious of Phactical Physiological Chemistry have never been a one-man job. The success and prestige of the book has been due in no small measure to the fine cooperation shown by many teachers and investigators in the medical and biochemical fields. A list of those who have participated in some degree on one or more editions would read like a combination of Wilo's Wito and Wilo Was Wilo in these professions.

In the claboration of this Edition the authors feel signally bonoted to have employed the cooperation of many distinguished scientists and educators To credit them individually with specific contributions would fail to do justice to those whose advice and assistance is reflected in several places, if not generally, throughout the book.

In consequence, deep grafitude is expressed here on behalf of my associated co-authors and myself for major contributions and assistance of the collaborators whose names and affiliations are listed immediately preceding this Preface. Without the splendid and gracious help of these eminent thou turnsts and physicians it would have been infinitely more difficult to produce what we behave is the finest of a long series of revisions.

Special arknowledgment and thanks are due the following eminent experts and teachers whose generous assistance; both solicited and unsolicited, while not quite as extensive as that of the aforementioned col-

Preface \1

laborators was none the less welcome Dr Zoe E Anderson, Director of the National Dairy Council's Research and Nutrition Service, Dr A K Balls, Professor of Enzyme Chemistry, Purdue University, Dr Albert L Chaney, Director of the Albert L Chaney Chemical Laboratory, Glendale, Cal , Dr L C Craig, Member, Rockefeller Institute for Medical Research, Dr Alexander L Dounce, Assistant Professor of Biochemistry. University of Rochester School of Medicine and Dentistry, Dr Theodore E Friedemann, Scientific Director, Medical Nutrition Laboratory, Pitzsimons Army Hospital, Denier, Col , Dr Linus Pauling, Professor of Chemistry, California Institute of Technology, Dr Kurt G Stern. Adjunct Professor of Biochemistry, Polytechnie Institute of Brooklyn, Dr Henry Tauber, Associate Professor of Experimental Medicine, School of Public Health, University of North Carolina, Dr Osear Touster. Associate Professor of Biochemistry, Vanderbilt University School of Medicine, Dr Everett C Cogbill and Mr Richard M Rush of the University of Virginia, and Dr M A Derow of Boston University

Once again it is my pleasure to pay tribute to the skill and untiring efforts of my associates, Dr Bernard L Oser, whose more than quarter-eentury association with this book is well known to its friends, and Dr William H Summerson, whose long experience as a teacher and investigator at Cornell Medical College preceding our association, eminently qualified bim for the important role be has assumed as a co-author

For their assistance in matters editorial the authors are particularly indebted to Mrs. Eumice Stevens, editor-in-chief of Blakiston, and her able and conscientious associate editor, Mr. Barney Pisha Their efforts in standardizing style and typography are reflected in the enfanced appearance of this completely reset edition. In this connection reference may he made to questions of spelling and nomenclature where uniformity of practice in the scientific literature has yet to be achieved. Since the American Chemical Society, more than any other single organization has been concerned with standardization in this field, the recommendations of that organization have been adopted despite the risk, in some instances, of jarring the sensibilities of a few readers (and occasionally of a co-author). While appreciation is expressed to the publisher's and printer's stuffs for their careful efforts in proofreading the authors assume full responsibility for such errors and oversights as will inevitably be discovered by sharp-eyed readers.

Appropriate acknowledgment has been made through this edition to the numerous authors publishers and instrument and apparities companies who so gracously grinted permission for the use of illustrations, tables, or other copyrighted material. Appreciation is expressed here for these courtesies and, in the unlikely event that acknowledgment has been useful to specific places where such material has been used, the authors claim human fruity and apologize for these oversights.

Finally, the authors are deeply appreciative of the patience and cooperation of the Maple Press whose fine craftsmaniship has made possible the production of this attractive volume.

I trust I will be pardoned for concluding this Preface with a short

personal note

The manuscript of the First Edition of this book was drafted high up in the old Hare I aboratory of the Medical School of the University of Pennsylvania, amid the stimulating fragrance of the Department of Anatomy It served as the basis for a course of instruction even before the edition was in print Now, a half century later, and thanks to its friendly reception by the profession the Thirteenth Edition has rolled off the press

Imong those to whom I was indebted for its genesis first place must go to the late Professor Lafayctte B Mendel for it was he who generated the spark of inspiration. The value of Dr. Mendel's suggestions and entities of the manuscript of several editions was great indeed. His wonderful letters written in long hand with purple ink and eramined full of ideas references, and suggestions were of incalculable aid. These missives were never less than six pages long and often twice as long. The lirst Edition was thus, quite naturally, based on the courses as given at Yale University, with minor features as adopted at Columbia University (College of Physicians and Surgeons) where I passed my first teaching assignment (1901–1903) under Professor William J. Gies. Dr. Mendel's interest in the book never flagged. It is therefore fitting and proper that to Dr. I fafayette B. Mendel guide, philosopher, and friend of many an outstanding book hemist as he was of the senior author, should be dedicated thus "Golden Anniversary." Edition of Priverical Physiological Ciri busying

April 1954 Viami Brach, Florios PHILIP B HAWK

Preface to the First Edition

The plan followed in the presentation of the subject of this volume is rather different, so far as the author is aware, from that set forth in any similar volume. This plan, however, he feels to be a logical one and has followed it with satisfactory results during a period of three years in his own classes at the University of Pennsylvania. The main point in which the plan of the author differs from those previously proposed is in the treatment of the food stuffs and their digestion.

In Chapter IV the "Decomposition Products of Proteids" has been treated although it is impracticable to include the study of this topic in the ordinary course in practical physiological chemistry. For the specimens of the decomposition products, the crystalline forms of which are reproduced by original drawings or by microphotographs, the author is

indebted to Dr Thomas B Osborne, of New Haven Conn

Because of the increasing importance attached to the examination of feces for purposes of drignosis the author has devoted a chapter to this subject He feels that a careful study of this topic deserves to be included in the courses in practical physiological chemistry, of medical schools in particular The subject of solid tissues (Chapters XIII, XIV and XV) has also been somewhat more fully treated than has generally been customary in books of this character

The author is deeply indebted to Professor Lafayette B Mendel of Yale University, for his careful criticism of the manuscript and to Professor John Marshall, of the University of Pennsylvania for his prinstaking revision of the proof He also wishes to express his gratitude to Dr David L Edsall for his criticism of the chinical portion of the volume, to Dr Otto Folin for suggestions regarding several of his quantitative methods, and to Mr John T Thomson for assistance in proof reading

For the micro-photographs of on haemoglobin and haemin reproduced in Chapter XI the author is indebted to Professor E T Reichert, of the University of Pennsylvania, who in collaboration with Professor A P Brown, of the University of Pennsylvania, is making a very extended investigation into the crystalline forms of biochemic substances. The micro-photograph of aliantofu was kindly furnished by Professor Mendel The author is also indebted for suggestions and assistance received from the lectures and published writings of numerous authors and investigators.

The original drawings of the volume were made by Mr. Louis Schmidt whose eminently satisfactory efforts are highly appreciated by the author

PHILIP B HAWK

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Physicochemical Principles

Living matter differs from nonliving in its possession of certain characteristic properties such as growth, reproduction respiration, and motion. The science of physiological chemistry deals with the application of chemical and physicochemical principles and methods to the study of these phenomena. In the early days of the science this meant the analysis of foods entering the organism and of evereta leaving it it involved the study of the composition of the various tissues and organs of the body, the blood, the digestive secretions etc. In this way a great deal of information has been collected concerning the composition of hving matter and the fate of the substruces that are necessary for continuance of life and growth. Although far from complete these researches have progressed to the point where we now possess a fairly comprehensive picture of the gross changes that take place in protoplasm.

The experimental methods used, however, for the most part involved the destruction of the living cell. In recent years the emphasis has been placed on the mechanisms concerned in the reactions of the living protoplasm itself. Since protoplasm is largely water, this requires a study of the nature of solutions and the complex behavior of mixtures of electrolytes. Since the physical basis of protoplasm is colloidal in character, a study of the peculiar structure and properties of colloidal solutions is also necessary. Some of the more important physicochemical principles that are finding wide and fruitful application in the study of life phenomena are

discussed briefly in the following pages

THE COLLOIDAL STATE

True and Colloldal Solutions. Thomas Graham in 1861 classified all substances into two groups crystalloids and colloids, depending upon their ability to diffuse through membranes such as purchment According to Graham crystalloids readily passed through parchment membranes while colloids did not. We now recognize that matter cannot be classified in this minner since many typical colloids, such as certain proteins, are crystallizable, and practically all crystalloids may, under proper conditions, by brought into the colloidal state.

According to modern concepts, colloidal solutions, instead of being solutions of particular types of matter are solutions with a characteristic kind of structure. Substances such as glucose or sodium chloride, which form true solutions in water, disintegrate, when dissolved, into individual molecules or ions which are less than 1 mg (1 millionth mm) in diameter.

The smallest particle that can be seen with a high power light microscope has a diameter of about 200 mm By means of an electron microscope parti cles with a diameter as small as 10 mm are re idily made visible. When the particles of solute are larger than 200 mu they are said to be in suspension, on standing such particles will gradually separate out When however the solute is dispersed into particles which are intermediate in size between ordinary molecules such as exist in true solutions and the course part cles found in suspensions it is said to be in the colloidal state and solutions containing particles of that size are known as colloidal solutions or sols Sols which have become jellylike are called gels

Colloidal solutions true solutions and suspensions thus differ from each other fundamentally only in the size of the particles of solute (the disperse phase) dispersed in the solient (the dispersion medium) Because of the dimensions of the disperse phase colloidal solutions exhibit certain characteristic and unique properties (to be discussed in later sections) which confer upon them their great importance in the structure of hving protoplasm This importance resides in the fact that protoplasm is con sidered to be a complex system containing many different crystalloidal and colloidal components. Although the structure and properties of this system are too complex to permit exact characterization in the present state of our knowledge we may gain an insight into these questions by a study of similar though very much simpler, colloidal systems such as are discussed below

Preparation of Colioidal Solutions The relationship between col loidal solutions true solutions and suspensions indicated above suggests two general methods by which colloidal solutions may be prepared. These methods are classified as (1) condensation and (2) dispersion methods depending upon whether the colloidal particles are formed by aggregation of individual molecules or by disintegration of coarse particles of matter

CONDENSATION METHODS The principles underlying the preparation of colloidal solutions by condensation methods are similar to those in volved in ordinary precipitation reactions. In both processes the solution is permitted to become supersaturated with respect to some particular substance Such supersaturated solutions in the presence of suitable con densation nuclei develop molecular aggregates which continue to in crease in size as long as any available material remains in solution. In precapitation reactions this process of growth continues until the particles become visible in a microscope or to the naked eve when they flocculate from solution By proper regulation of the experimental conditions which differ for different substances and procedures the growth of molecular aggregates may be checked when the particles attain the size charac teristic of the colloidal state thus forming colloidal solutions. Whether a particular reaction will lead to the formation of a colloidal solution or a visible precipitate depends therefore entirely upon the conditions under which the experiment is carried out Von Weimarn who studied this question very extensively showed that by merely varying the concentra tions of Ba(CNS); and MnSO, from \/20 000 to 7 \ the form of the BaSO, precipitated could be made to vary from large crystals to a colloidal cel The colloidal state as indicated in the following scheme is merely an intermediate stage between coarse precipitates and true solutions which may be approached, under proper conditions, from either direction

Condensation Methods→
True Solutions

Molecules and Ions Diameter Less than 1 mµ Collordal Solutions
Molecular Aggregates
Diameter 1 mµ to
200 mµ

Suspensions
Molecular Aggregates
Diameter, Greater
than 200 m

-- Dispersion Methods

Most of the morganic colloids may be formed by condensation methods involving such reactions as reduction, oxidation, hydrolysis, and double decomposition Thus if a dilute solution of gold chloride is treated with formaldchyde under proper conditions, the gold ions are reduced to atoms of gold which then aggregate into particles of colloidal size Practically all of the metals yield colloidal solutions under similar conditions. Boiling a very dilute solution of ferric chloride results in hydrolysis of this salt with the formation of a colloidal solution of ferrie hydroxide Similar solutions may be obtained by using salts of Cr. Al, or Sn A dilute solution of arsonious oxide in water, when treated with hydrogen sulfide, undergoes double decomposition with the formation of colloidal arsenious sulfide. In the preparation of colloidal solutions by these methods it is essential that the reactions used do not lead to the formation of soluble, strong electrolytes This condition is important because colloidal solutions of the type discussed here are extremely sensitive to small amounts of electrolytes, which cause aggregation of the colloidal particles into larger particles which precipitate out of solution

Dispersion Methods The dispersion methods for preparing colloidal solutions involve, as indicated above, the subdivision of coarse material into particles of colloidal size under conditions which will prevent coalescence of those particles. Many substances may be reduced to approvimately colloidal size by grinding in a colloid mill, which consists essentially of two flat metal plates placed almost in contact and rotated at very high speeds in opposite directions Colloidal solutions of most of the metals may be prepared by producing an electric are between electrodes of the metal held under water or some other suitable liquid. It is quite probable that this method involves some condensation of metal vapors as well as disintegration of the metal itself. Solutions prepared by either of these methods will gradually flocculate unless some stabilizing agents are added to prevent coalescence of the partieles. For this purpose use is made of certain substances, either electrolytes or other colloids, called peptizing agents. The electrolytes usually act hy emferring an electrical charge on the colloidal particle, which is an essential condition for stability of colloidal solutions of this class Peptizing colloids, also called protective colloids, apparently form a film about the individual particles, which converts them into particles resembling the protective agent in stability and other properties

General Properties of Colloidal Solutions. It has already been pointed out that the essential difference between colloidal and true solutions lies in the size of the particles of solute dispersed in the solvent

Although the particles in a colloidal solution are too small to be retained by ordinary filter paper, they are too large to pass through such membranes as collodion parehment or cellophane, which are permeable to most substances in true solution. This inability of the colloidal particles to diffuse through certain membranes is made use of in the process known as dialysis, by which colloidal solutions may be freed from noncolloidal impurities. The solution is placed in a suitable dialyzing hag and is suspended in a large volume of distilled water which is changed at frequent intervals until the liquid inside the bag no longer gives a test for the particular substance to be removed. This process of dialysis is used extensively in the preparation of salt-free solutions of such blodgical colloids as the proteins Last traces of electrolytes may be removed by the process known as electroal alysis, in which an electrical current is passed through the solution in a suitably designed apparatus.

If a solution containing colloidal particles is forced through a suitable membrane by pressure, the membrane becomes an ultrafilter and the colloidal particles are separated from the solution. This process is called ultrafiltration. By this means it is possible to separate the crystalloidal and colloidal components of such a fluid as blood plasma, for example Furthermore it is possible to prepare membranes containing porces of almost any desired size so that colloidal particles of any particular size may be separated from smaller particles as well as from material in true solution.

Another widely used method for separating colloidal particles of different sizes is hased upon the use of the high speed centrifuge or ultracentrifune This apparatus originally developed by Svedherg for the study of particle size in suspensions of morganic colloids, has found extended application to the problems of protein and virus chemistry by Svedberg and many others, until today it is a fundamental research instrument in these fields With this instrument a sedimenting force up to several hundred thousand times the force of gravity can be applied to particles in colloidal suspension Naturally the heavier particles will settle out of the solution under these conditions faster than the lighter or smaller particles, and the actual separation and isolation of individual colloidal substances has been achieved in this way Furthermore by the use of suitable optical devices it is possible to follow the rate of settling in the various types in colloidal particles which may be present and thus establish not only whether the disperse phase consists of colloidal particles of several different sizes (heterogeneous) or of only one size (homogeneous), but also the average size of the particles themselves. If it can be shown that the colloidal particle is a single molecule of the substance, the particle size then becomes a measure of the molecular weight of the substance. This method of approach has found particular application in the field of protein chemistry

The sedimentation of a particle under stated conditions may be char acterized by the sedimentation constant s defined as

where dx/dt is the rate of settling of the particle per unit time at n distance x from the center of rotation of the centrifuge, and ω is the angular velocity in radians per second. Since the numerical value of s is usually very low—around 1×10^{-13} for proteins—sedimentation constants are commonly expressed in terms of a unit, the Svedberg, S, which equals s multiplied by 1×10^{13} .

The relation between the sedimentation constant and the molecular weight under ideal conditions is given by the following expression:

$$M = s \cdot \frac{RT}{D(1 - \bar{v}d_m)}$$

where M is molecular weight, R is the gas constant, T the absolute temperature, D the diffusion constant, \bar{v} the partial specific volume of the substance (the volume occupied by one gram of substance at infinite dilution in the solvent) and d_m the solvent density. This equation requires separate determination of the diffusion constant D; errors in D will be reflected in the value for the molecular weight. Nevertheless the equation, suitably corrected for deviations from ideal conditions, has been widely used. Examples of the molecular weights of proteins determined by sedimentation velocity will be found in Chapter 5.

An alternative procedure for the determination of particle size or molecular weight in the ultracentrifuge is based upon the attainment of a sedimentation equilibrium. In this the rate of settling of the particles is established at such a point that it exactly equals the opposing force of diffusion, which tends to redistribute the solute throughout the solvent. Under these conditions the molecular weight is obtained by means of the equation:

$$M = \frac{2RT \ln \frac{C_2}{C_1}}{\omega^2 (1 - \bar{v}d_m)(x_2^2 - x_1^2)}$$

where C_1 and C_2 are the concentrations at distances x_1 and x_2 , respectively, from the center of rotation. (The symbol "In" stands for the natural logarithm.) The value of the diffusion constant is not required in this method, and this is an advantage. On the other hand, precise establishment of sedimentation equilibrium requires eareful control over sedimentation rate for relatively long periods of time. In general, molecular weights determined by sedimentation equilibrium differ but slightly from those obtained by sedimentation velocity.

By noting and measuring deviations from the equations just presented, which are based upon spherical particles obeying the gas laws, information concerning not only the size but also the shape of particles may be obtained. In the case of proteins, variation in the value of the sedimentation constant with concentration has been interpreted in terms of equilibrium between monomeric and dimeric forms of the protein molecule.

Since the particles in a colloidal solution consist either of very large molecules or of aggregates of large numbers of individual molecules, it follows that a colloidal solution will contain only a minute fraction of the number of particles present in a true solution of the same concentration Such physicochemical properties of solutions as vapor pressure and osmotic pressure depend almost entirely upon the number of particles in solution, the chemical nature of these particles being immaterial Colloidal solutions, with their comparatively small number of particles, therefore exhibit only very small osmotic pressures, and their vapor pressures, as well as related boiling and freezing points, are practically the same as those of the pure dispersion medium

If a concentrated beam of light is passed through a colloidal solution and viewed at right angles, its path through the solution is plainly visible as a milky turbidity. True solutions under similar conditions appear optically void This phenomenon, known as the Tyndall effect, is due to the reflection and scattering of the light by the particles in solution and is similar to that observed when a beam of sunlight passes through the

dust-laden atmosphere of a darkened room

Siedentopf and Zeigmondy, in 1903, developed the ultramicroscope, in which a very fine, intense beam of light is focused in a colloidal solution and its path observed with a microscope placed at right angles to the beam The true nature of the Tyndall effect becomes apparent when colloidal solutions are viewed with the ultramicroscope It is then observed that the Tyndall light is composed of individual points of light, each point representing light reflected by an individual particle Colloidal solutions are thus shown to consist of discrete particles suspended in the dispersion medium. When viewed through the ultramicroscope the particles in a colloidal solution are seen to be in a continual state of violent vibratory motion This phenomenon is known as the Brownian movement and is due to bombardment of the particles of disperse phase by the molecules of the solvent

Colloidal particles ordinarily carry electrical charges which are considered to be distributed over the surface of the entire particle Because of the presence of these charges the partieles in a colloidal solution migrate toward one of the poles in an electrical field, the process being

known as cataphoresis or, more commonly, electrophoresis

The distance x through which a particular colloidal particle moves under the influence of an electric current is proportional to the time t and to the potential gradient E at the spot where the particle is E in turn depends upon the current 1, the conductivity of the solution A, and the cross-sectional area q of the solution at this point. The proportionality constant is known as the electrophoretic mobility # Hence $\mu = x/tE = x/t \times q\lambda/t$ The dimensions of μ are therefore square centimeters per volt per second. The distance x is usually measured by the so-called moving houndary method. In the version of this procedure developed by Tisehus and widely used, the colloidal solution is brought into contact with the solvent alone by a process of layering that uses a sectioned U-tube (see Fig 105, p 461) to produce a sharp boundary between the colloid-containing solution and the solvent alone, although the two solutions are continuous through the solvent itself. An electric current passing across this boundary will carry the colloidal particles into the solvent or away from it, depending upon the polarity of the current and the charge on the particles If all the particles are similar with respect to charge, mass, and shape, they will move as a unit, so that in effect the boundary itself will move and will remain sharp. Optical or other methods for measuring the movement of the boundary will then establish the rate of movement of the colloidal material. If the colloidal solution is composed of various types of particles with differing electrophoretic mobilities, several boundaries or zones will develop, moving at different rates. In this way a mixture of colloidal substances may be characterized, and under suitable conditions the components may actually be separated from one another.

A simplified form of electrophoresis apparatus suitable for clinical and other uses employs a strip of filter paper (paper electrophoresis) saturated with solvent (electrolyte) and attached to two electrodes. The colloidal solution is deposited on the filter paper at one spot, the filter-paper matrix tending to localize the added material. Under the inflinence of an electric current the colloidal particles present will migrate along the paper strip in bands in much the same manner as that described above for the Tiselius procedure, and these bands may be visualized by suitable procedures (see Fig. 57, p. 184).

Like the ultracentrifuge mentioned above, electrophoresis in the study of colloidal systems has found application not only in isolating individual colloidal substances from mixtures but also in characterizing the components of complex natural colloidal solutions such as the proteins of

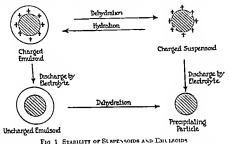
blood plasma

Cinssification of Colloidal Solutions. Colloidal solutions are classified into (1) suspensoids, or lyophobic colloids and (2) emulsoids, or lyopholic colloids. The suspensoids include such colloidal solutions as those formed by the metals, inorganie salts, etc., and their preparation usually requires special methods such as the dispersion and condensation methods described above. They are called lyophobic colloids because there is no affinity between the particles of disperse phase and the dispersion medium. In many of their physical properties, such as viscosity, suspensoids differ only slightly from the pure dispersion medium. The particles of a suspensiod carry a definite electrical ebarge which may be changed only by very special methods. Suspensoids are focculated by very small quantities of electrolytes, and when so precipitated cannot ordinarily be brought back into colloidal solution. The precipitation of suspensoids is therefore irreversible. When viewed with the ultrameroscope suspensoids are seen to contain well-differentiated particles in vigorous Brownian movement.

The cmulsoids, since they include the proteins and the higher carbohydrates, are of much greater biological importance and interest than the suspensoids. They are called lyophilic (or hydrophilic when the dispersion medium is water) colloids because they have a high affinity for the dispersion medium. The viscosity of emulsoids is usually much higher than that of the dispersion medium. The particles earry electrical charges which, in the case of proteins, for example, may be changed in sign or magnitude by such simple measures as changing the acidity or alkalimity of the solution. The emulsoids require large amounts of electroly tes for their precipitation and when precipitated may issually be brought back into colloidal solu-

tion by the addition of fresh solvent. The precipitation of emulsoids is therefore reversible

Emulsoids and suspensoids thus exhibit essential differences in their behavior toward the solvent and in their sensitivity to small amounts of electrolytes The stability of suspensoids depends entirely upon the charges on the particles, which permit them to repel each other and thus prevent aggregation into coarser particles. If the charges on the suspensold particles are neutralized, or reduced below a certain critical value, the particles coalesce and precipitate The sensitivity of suspensoids to small amounts of electrolytes is therefore due to the fact that electrolytes neutralize or reduce the charges on the particles and thereby remove the



force which is ordinarily sufficient to prevent aggregation and floeculation In this precipitating action of electrolytes on suspensoids, the active ion is the one whose charge is opposite in sign to that carried by the suspensoid

On the other hand, emulsoids possess two stability factors-charge and hydration-either of which is capable of preventing the aggregation and flocculation of the colloid Neutralization of the charges on the cmulsoid converts it into a neutral, or recelectric, colloid which is perfectly stable as long as the particles remain hydrated Dehydration of a charged, electrolyte-free emulsoid (e g by the addition of a dehydrating agent like alcohol) converts it into a suspensoid exhibiting the characteristic sensitivity toward electrolytes The precipitating effect of large amounts of such salts as ammonium sulfate or sodium chloride on emulsoids is due to the fact that saturated solutions of these salts act also as delaydrating agents, thereby discharging and dehydrating the particles at the same time Fig 1, from Kruyt illustrates these differences in stability between emulsoids and suspensoids

Loder changing conditions of temperature and concentrations of disperse phase hydrogen ions and electrolytes emulsoids have the property of imbibing large quantities of water and setting into semirigid gels. These gels are thought to possess a definite structure, consisting in some cases of a network of disperse phase, or hydrated material, enclosing some of the dispersion medium. On standing for a time most gels gradually contract, extruding a portion of the dispersion medium. This process is known as syncress. The properties of imbibition and gel formation exhibited by emulsoids probably are responsible for the characteristic physical form of protoplasm. The swelling and shrinking exhibited by protoplasm are due probably to the hydration and dehydration of its colloids, presumably governed by the same laws that operate in the case of the simpler systems.

Colloidal particles possess appreciable surfaces which permit such phenomena as surface tension (qv) to come into play at the interfaces between the particles and the dispersion medium. The magnitude of these surfaces becomes evident when we reflect that if a 1-cm cube is subdivided into cubes with 10 mm edges, the surface area is increased from 6 sq cm to 600 sq m Substances such as most emulsoid colloids (soaps, proteins, etc) which decrease surface tension tend to accumulate at surfaces, whereas other substances such as sodium chloride which increase surface tension tend to be less concentrated there. This increase in the concentration of a substance at a surface is known as odsorption. In the ease of colloidal solutions, adsorption thus involves an increase in concentration in the boundary layer between dispersed phase and dispersion medium The mechanism of adsorption is obscure, it has been suggested that the adsorbent enters into a loose combination with the material adsorbed by means of latent valences of atoms in the surface layers. In many cases (such as adsorption of many different kinds of gases, liquids and even solids by such suspensions as charcoal), it is difficult to understand how any kind of chemical combination takes place. When electrolytes are adsorbed by the charged particles in colloidal solutions it is the ion with opposite charge which is adsorbed

Adsorption probably plays a very important role in many of the reactions taking place in living protoplasm, since in this manner substances ordinarily present in low concentration may have their effective concentrations increased tremendously by being accumulated at boundary surfaces. Concentration by adsorption also finds use in the laboratory in the isolation from very dilute solution of such natural products as enzymes, vitamins, bormones, etc. The adsorbed material may be released (cluted) from the surface of the solid adsorbent (e.g., charcoal, metallic hydroxides, fuller's earth) by changing the activity or surface tension of the solvent. In the process of chromatographic adsorption the solid adsorbent is packed into a tall glass column down which the solution containing adsorbable material is allowed to flow slowly. This procedure is discussed in detail in the section on obromatography (see p. 14)

The studies of Harkins and of Langmur indicate that substances aceumulate at boundary surfaces in a definite pattern. Thus molecules having both polar and nonpolar groups orient themselves at oil-water interfaces in such a fashion that the polar groups dissolve in the water while the nonpolar groups dissolve in the oil. Oleie acid, for example, orients itself at oil-water interfaces so that the polar carboxyl group is directed into the water, which it more closely resembles in general structure than does the nonpolar alkyl group, which is directed into the oil. Since protoplasm contains both water and lipide materials it is quite probable that some of the cell constituents are oriented in a similar fashion within the cell. It is also probable that the internal structure of the cell is based upon orientation of various types of molecules within the cell.

EXPERIMENTS ON COLLOIDAL SOLUTIONS

A METHODS FOR PREPARATION OF COLLOIDAL SOLUTIONS

 Preparation of Colloidat Solutions of Prussian Blue and of Arsenious Suffide by Double Decomposition. To 10 ml. of 0.02 N potassium ferrocy ande in a beaker add with mixing 10 ml. of 0.02 N ferric chloride. Dilute a portion of the mixture and note that there is no precipitate. This colloid is negatively.

Mix 50 ml. of 1 per cent arsenious oxide solution (made by boiling the oxide with water) with 50 ml. of saturated solution of hydrogen sulfide. Heat to holling (in the hood), filter and coof, This colloid is electronegative also.

- Preparation of Colloidal Ferric Hydroxide by Hydrolysis. To 290 ml. of holling water add 1 ml. of 33 per cent ferric chloride solution, or about 0.3 g. of solid ferric chloride. Note the beautiful reddish-brown color. This is a positively charged colloid.
 - 3. Preparation of a Gold Sol by Reduction. To 100 ml. of pure redistilled water add 1 ml. of 1 per cent gold chioride solution and 5 ml. of 0.5 per cent potassium carbonate solution. Heat to boiling, remove from the flame, and add two drops of 20 per cent formaidehyde solution. If necessary, heat again and add one more drop. Observe the changes in color as the particles of metalic gold become larger and larger. Most metallic colloids are electronegative.
 - 4. Preparation of Colloidal Platinum by Electrical Method. Connect two short platinum wires (1 to 3 mm. cross-section) with stout copper wires and pass through pieces of glass tubing to serve as handles. Connect both wires with the terminals of a 110-V. d.c. lighting circuit with a lamp bank of five 1800-watt lamps in parallel (to cut down the current to about 5 amperes). Place pure distilled water in a crystallizing dish about 4 inches in diameter, add a very slight trace of bydrochloric acid, and immerse the two platinum tips. Bring the tips together and then separate them by about 1 to 2 mm. An arc develops. Maintain this for several minutes. Filter.
 - 5. Preparation of Emulsoid Solution, Prepare solutions of gelatin (5 per cent), agar (1 per cent), starch (2 per cent), and soap (2 per cent) by allowing the dry colloid to become thoroughly soaked in the required amount of water in a beaker at room temperature, followed by immersion of the beaker in a boiling, water bath, attring until solution is complete. Cool the solutions and note that gels are formed. Heat again in holling, water and note that the gels liquety. On standing for some time the gels should show syncresis.

B. General Properties of Colloidal Solutions

 Diffusion Through Membranes (Dialysis). Prepare a dialyzing bag from seamless cellulose dialyzer tubing as follows: Cut a strip about 15 cm. long

Obtainable from most isboratory supply houses in the form of rolls of varying length and ridth. If not sysilable, collodion bases may be used. Pour 10 to 20 ml of collodion solution.

from the roll of flat tubing, wet the strip thoroughly with water, and separate the sides to form a tube. (This operation is facilitated by holding the strip vertically in a stream of running water.) The one end of the tube tightly with string to form a bag. Fill the bag with colioidal Prussian blue or ferric hydroxide solution (prepared as described above), tie loosely at the top with string, and suspend in a beaker of water. Observe at intervals. Do you detect any diffusion of the colloid through the membrane? Test the water in the beaker for chloride. Is any substance diffusing through the membrane?

- 2. Diffusion Through Gels. Place 5 ml. of 5 per cent gelatin in each of two test tubes. Cool until the gelatin solidifies. Pour copper sulfate over the gel in one tube and Prussian blue solution over the other. Let stand. Note the degree of diffusion of the blue color in the two cases. The experiment may be repeated with eosin and methylene blue, the former a noncolioidal and the latter a colloidal dye.
- 3. Mutual Precipitation of Colloids. Mix equal volumes of solution of the negative colloid arsenious sulfide and the positive colloid ferric hydroxide. Explain. Can you write the equation for any reaction which occurred?
- 4. Precipitation of Colloids by Salts. To a solution of Prussian blue, ferrle hydrotide, or other suspensoid, add a few ml. of 10 per cent sodium chloride solution, and allow to stand. Treat a solution of an emuisoid such as gelatin or starch in the same way. Which are precipitated by the electrolyte? Add an excess of water to the precipitate. Is the colloid reversible? If the emulsoid is not precipitated saturate the solution with solid magnesium sulfate. To some of the precipitate add an excess of water and boil. Is the emulsoid reversible?
 - 5. Demonstration of a Protective Colloid. To a mixture of three drops of concentrated nitric acid and 5 mi. of 0.05 N sliver nitrate add 5 ml. of 0.05 N sodium chioride. Note the curdy precipitate. Repeat, but add 1 ml. of gelatin solution to the silver nitrate and to the sodium chioride solutions before mixing.

7. Viscosity (Internal Friction) of Colloidal Solution. Fill a 10-ml. pipet with water, suspend it in a vertical position, and allow it to drain. Count the number of seconds required for emptying. Repeat, using a solution of Prussian blue or ferric hydroxide. Repeat again, using a 2 per cent solution of gelatin, ilow does the viscosity of the suspensold compare with that of the emulsoid and with that of pure water? A more accurate form of this apparatus is a viscosimeter of the Ostwald type (see Exp. 8). A solution of gelatin shows its lowest viscosity (most rapid emptying of the viscosimeter) at its isoelectric point (about pil 4.7).

8. Estimation of Viscosity with Ostuald Viscosimeter. The apparatus illustrated in Fig. 2 is used and is kept in a constant-temperature water bath dur-

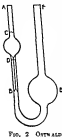


FIG. 2 ORTWAL VISCOSIMETER

ing the determination. Test with water first. By means of a pipet introduce a suitable exact amount (usually 5 mi.) of water into bulb E. Blow at F and force the liquid abore mark G. The fluid column should still reach to huib E. Allow it to flow back and note the time with a stop watch for the meniscus to pass from G to D. Repeat until constant values are obtained. Repeat with gelatin or some other colloidal solution, if water requires 60 seconds and gelatin solution 120 seconds, the relative viscosity of the gelatin is 2 0.

9. Adsorption by Charcool. Caramelize a small amount of sugar hy heating it in a crucible. Dissolve in 100 ml. ol water, add 5 g. of animal charcoal, and boll for five minutes. Filter. The filtrate should be coloriess. Charcoal is used for clarification in the commercial refining of sugar and in many slimitar processes. Repeat the experiment, using a dilute solution of crystal violet or other dye instead of the caramelized sugar solution. After the filter bas drained, pour alcohol or acctone over the moist char-

coal. Does the dye enter into a firm combination with the charcoal? Acetone and alcohol are more strongly adsorbed than crystal violet and tend to displace it from the charcoal. What is this displacement process called?

10. Electrophoresis of Colloids. Champ in an upright position a glass U-tube (one with limbs ahout 15 cm. long and 2 cm. in diameter is satisfactory). Italf fill the tube with a satisfate subvived colloids solution (knewnors suffactory). For its satisfactory) and carefully add distilled water to each limb to within about 2 cm. of the top. With suitable care a sharp boundary between the water and the colloids solution is obtained. Place a platinum electrode in contact with the water in each limb, and connect each electrode to a terminal of known polarity from a source of 110 v. d.c. Note (1) the level of the colloid-water boundary in each limb and (2) the polarity of the electrodes, and turn on the current. Allow it to run until definite movement of the boundary has taken place. What does this experiment prove with regard to the sign of charge on the colloidal particles? Repeat with a colloid of oposite charge.

DONNAN EQUILIBRIUM

The British physicist Donnan has made a contribution of the utmost importance in physiology to our knowledge of the influence of ions in colloidal systems. Let us suppose a membrane (indicated below by a vertical ine) to be impermeable to the anion R of a salt NaR in solution on one side of the membrane but permeable to all other ions and salts involved. Let us further suppose that we have at the beginning of the experiment NaR on one side of the membrane and NaCl on the other. (See diagram at left, below.) NaCl will diffuse from 2 to 1 until at equilibrium conditions will be as indicated on the right.

Now from thermodynamic considerations it can be calculated, and by actual determination it can be shown, that at equilibrium the product of the concentrations of a pair of diffusible ions (in this case Na⁺ and Cl⁻) on one side of the membrane will be the same as the product of the concentrations of the same ions on the other side of the membrane. That is,²

$$[Na+]_1[Cl-]_1 = [Na+]_2[Cl-]_2$$

The concentration of the cation Na⁺ in (1) must be equal to the sum of anions present [R⁻ + Cl⁻] in order to maintain electrical neutrality, while on the other side the concentration of Na⁺ is the same as that of the anion Cl⁻. It follows that, in order for the products just mentioned to be equal, the concentration of sodium ions [Na⁺] on the left must be greater than on the right and the reverse must be true for the chlorine ions. That is,

$$[Na^{+}]_{1} > [Na^{+}]_{2}$$
 and $[Cl^{-}]_{1} < [Cl^{-}]_{2}$

Thus it is possible to account for the existence of different concentrations of diffusible ions on the two sides of a semipermeable membrane, the essential condition being the existence of a nondiffusible ion on one side of a membrane which is permeable to the other ions.

This difference in concentration of diffusible ions leads to a difference of potential (electrical pressure, measured in volts) on the two sides of the membrane, and may account for certain electrical phenomena of living matter. The Donnan equilibrium has also been associated with the explanation for a variety of physiological phenomena, such as the formation of gastric juice and other secretions, certain aspects of absorption and elimination, and the differences in concentration of diffusible ions between the blood plasma and ecrebrospinal fluid and between the red cells and plasma. In this connection it is well to remember that quantitatively the Donnan equilibrium is based upon ion activity rather than concentration, which was used in the above illustration for purposes of simplicity, and it is not always possible to define precisely the relation between these two properties, particularly in physiological fluids.

EXPERIMENTS ON DONNAN EQUILIBRIUM

- I Demonstration of Donnan Equilibrium Using a Membrane introduce 59 ml of a 2 per cent gelatin solution into one 100 ml beaker and 50 ml of distilled water into another Add 1 ml of a 64 per cent this mol blue solution to each To the water add 0 i N hydrochloric acid solution cautiously, drop by drop, until the indicator has a color intermediate between yellow and pink (pil about 2). Add the acid to the gelatin solution until a similar shade of color is ohtained (several ml may be required). The two solutions now have the same hydrogen ion concentration. Transfer the gelatin solution to a dialyzing bag and suspend it in the beaker containing the water solution. Let it stand. The gelatin solution will gradually, turn yellow (decreased acidity) and the outside solution will turn pink, (increased acidity), indicating a diffusion of hydrogen ions through the membrane. Do any other ions diffuse? What is the sign of charge on the nondiffusible gelatin ion in this experiment? Explain the results in terms of the Donnan equilibrium.
 - 2 Demonstration of Donnan Equilibrium without Use of a Membrone Prepare 190 ml of a 5 per cent solution of gelatin Add 1 ml of a 0.5 per cent solution of brom thymol blue Add 6 5 per cent % A011 drop by drop until a color intermediate between yellow and blue (pil about 7) is obtained Poul it into a 2-0 ml beaker and allow it to solldify Over the gel pour 100 ml o' distilled water containing the same amount of indicator and brought to this same pil Let it stand, and note the changes in the color of both gel and solution Explain is the presence of an extraneous membrane essential for the development of the Donnan equilibrium?

CHROMATOGRAPHY

Chromatography is the name originally applied by the Polish botamist Tswett in 1906 to a procedure for separating a mixture of different-colored pigments (chlorophylls and xanthophylls) from each other. Tswett found that if the mixture of pigments in petroleum ether as solvent was poured onto the top of a column of calcium earhonate firmly packed in a narrow glass tube, and allowed to drain down the column, the pigments were separated and appeared as colored zones along the column. By the addition of fresh solvent the zones could be further separated and the reparate pigments identified under suitable conditions. Tswett called such a column achromatogram and the process the chromatographic method.

As has happened with many other fundamental discoveries, the useful ness of Tswett's simple procedure was not generally appreciated until years later. At the present time chromatographic procedures of one kind or another are probably more widely used in hochemical research than any other single type of procedure. Tswett ascirbed the differences in the behavior of pigments placed on his columns to differences in adsorbability, the more strongly adsorbed chlorophylls for example displacing the weakly adsorbed xanthophylls and thus forcing the xanthophylls further along the column under the influence of the flowing solvent. There are however instances which are more readily explicable on some other hasis. For example, a mixture of amino acids placed on a column of starch or powdered cellulose may be chromatographed and separated by the use of certain solvents, but the order of separation of the various compounds

present parallels their distribution coefficients (see p 24) between the solvent and water Since starch or cellulose will take up appreciable amounts of water2 it is reasonable to assume that the amino acids dissolve in the water of the column and are extracted from it by the flowing solvent in proportion to their relative solubilities in the two phases (partition chromatography) In columns containing ion exchange resins (see below). it is believed that the functional groups of the resin react directly with the various substances placed on the column, but to a different degree Thus a resin containing acidic functional groups may be expected to react with a mixture of bases placed on the column in the same way that an acid in solution distributes itself between various bases present. In such a column the bases will be distributed along the column, the most reactive base at the top, and upon elution they will emerge from the column at different rates for similar reasons. There are therefore various explanations for the behavior of substances undergoing chromatographic separa tion, and the science is still far behind the art in this field Columns satisfactory for particular purposes are ordinarily found by trial, the explanation usually comes later

Suitable columns have been made from the various insoluble alkalineearth phosphates and oxides, starch and cellulose, and ion-exchange resins The dimensions of the column may vary, but in general a column resembles the common volumetric buret in ratio of length to width Every column has an inherent capacity for the particular substances heing fractionated on it, if this capacity is exceeded separation will be unsatisfactory If the various substances on the column can be released by solvent change and thorough washing a used column can be regenerated and used again The solvent may flow down the column by gravity or more commonly under a controlled pressure head. The rate of flow of solvent may vary in general, the more slowly the solvent flows, the more thoroughly equilibrium is reached and the potentialities of the column are realized. The solvent may be aqueous or nonaqueous as required, and may be changed during a separation to modify distribution along the column The wide variety of solvents available is one of the outstanding advantages of chromatography

The applicability of chromatography is not limited to colored compounds, thus the name is somewhat misleading. Any procedure which identifies a substance may be used to locate it in the column Fluorescence under ultraviolet light has been used, as has radioactivity, in cases in which the substance under study is labeled with a radioactive isotope. The column may also be extruded from the tube and cut into sections, and each section may be extracted and analyzed for the substance sought an improved isolution procedure now widely used is based upon the continuous passage of the pure solvent down the column, the effluent solvent being collected in separate fractions and the fractions analyzed for the material under study. The solvent may be varied during this process to promise the solvent may be varied during this process to

N + 1

earefully controlled conditions it is found that a particular compound may show sufficient reproducibility with regard to the fraction of the effluent in which it is found to afford a tentative basis for identification, this procedure has been used by Moore and Stein in their analysis of the amino acids present in protein hydrolyzates

Ion-exchange Resins. The use of ion exchange resins in chromatog raphy and for other purposes in biochemistry and medicine is of sufficient importance to justify brief consideration at this point Ion-exchange resins are highly insoluble synthetic polymers containing accessible (i.e., titratable) functional groups which are either acidic (—COOH,—SO₂H)

or basic (—NH₂ —N(CH₃); etc.) Resins with acidic functional groups such as Dower 50, Amberlite IR 100, and Permutit H, are known as cation exchange resins those with basic functional groups, such as Dower 1, Amberlite IR 4 and PermutitS are the anion-exchange resins

The action of resins is best understood by considering them to be poly functional acids or bases which happen to be insoluble. Thus if a solution containing an amino salt is placed on a column packed with a cation exchange resin containing replaceable hydrogen, the resin will take up the amine cation from the solution and replace it in solution by the hydrogen ion (hydrogen cycle) The amine may be subsequently displaced from the resin and washed out of the column by the use of a more acid solvent A mixture of hases on such a column may be released individually by the use of solvents of varying acidity and electrolyte content. In a similar manner, a cation-exchange resin whose bydrogen has been replaced hy sodium will function in a sodium cycle toward other cations such as calcium and magnesium A solution containing a calcium sali will emerge from such a column as a solution of the sodium salt the calcium remaining attached to the resin. The resin may be returned to its original state by treatment with a large excess of sodium ions. This is a principle used in the softening of hard water. An amon-exchange resin which has been saturated with chloride ion will function in a chloride cycle towards other amons a column containing such a resin will remove weak organic acids from solution as amons and replace them with chloride

The use of ion-exchange results is not confined to chromatography, but is acquiring increasing importance in other aspects of biochemistry and medicine. Examples of such uses include the use of results to decaledly blood and thus prevent blood clotting chinical use to control gastric acidity and the dictary absorption of electroly tes from the intestinal tract by the oral administration of suitable results and the demineralization of milk for special dictary purposes. Since it is now known that results can be "tailor made" to fit almost any desired purpose, such appear to be on the increase.

Paper Chromatography. A remarkable development in the field of chromatography which is perhaps inequalled in the field of biochemical procedures for its simplicity versatility and widespread application is the use of filter piper for chromatographic purposes by a procedure first described by Consden Gordon and Martin 4 The basic principle is very

Consider Gordon and Martin Biochem J 38 221 (1914)

simple A small drop of solution containing the mixture of compounds it is desired to separate is exaporated to dryness on a piece of filter paper, and a suitable solvent is allowed to flow slowly along the filter paper over this spot, either by gravity (descending) or capillanty (ascending). The substances in the initial spot are extracted by the flowing solvent and carried along the filter paper to an extent which is related to their distribution between the solvent and the water phase of the filter paper, as discussed previously. After the solvent has flowed for a suitable distance along the paper, the paper is removed and freed of excess solvent by quickly drying and suitable tests are applied to the filter paper to locate the various compounds under study. Under the proper conditions it is found that each substance present has been carried away from the initial spot to a characteristic extent and is localized in a relatively small area on the filter paper.

The simplest expression of the response of a particular compound to this procedure is the so-called $R_{\rm F}$ value, which is the ratio of the distance the compound moves along the paper to the distance covered by the solvent Thus if the solvent front is 120 cm. from the spot of application of the compound, and the compound is located 90 cm. from this spot, the $R_{\rm F}$ value is 9 0/120, or 0.75. The absolute value of $R_{\rm F}$ for a particular compound is of course dependent upon such factors as the nature of the solvent, the temperature, and the presence of other substances which influence the distribution of compound between the solvent and water. With suitable attention to these details, however, the $R_{\rm F}$ value for a pure substance is found to be sufficiently reproducible to serve as an approximate basis for identification.

An improved procedure is the so-called two dimensional chromatography. Here the spot containing the substances under study is dried in one corner of a square of filter paper. The solvent, running as before, curries the substances along one edge of the paper in proportion to their R_T values for that solvent. After the excess solvent is removed by drying the paper is turned 90° and a second solvent is allowed to flow across the paper at right angles to the direction of flow of the first solvent and in such a way as to carry, the various substances present from the edge of the paper to the middle regions. After the second solvent is removed by drying, the substances are located by suitable means. By the selection of suitable solvents it is possible using this procedure to separate and identify the components of even such complex mixtures as the various amino acids present in a protein hydrolyzate (see Trontispiece) using only small amounts (a few milligrams) of material and ordinary laboratory equipment

There are of course many refinements to the procedures described here By the careful use of the technique, the compounds separated are usually localized in a small and reproducible area of the filter paper. If the compound is detected at this spot by a color reaction, measurement of the color intensity may be used to determine the amount of material present. The area may also be cut out ind the compound extracted from the paper, thus essentially solating it in pure solution for confirmation of its chemical nature by other tests. The student will find many applications of paper chromatography to biochemical problems in this book, and in the bio-

chemical literature

EXPERIMENTS ON CHROMATOGRAPHY

1. One-dimensional Paper Chromatography.* For demonstration purposes solutions of amno acids will be chromatographed in this experiment. The amino acids will be located on the filter paper by reaction with ninhydrin to give a purple color. Solutions of glycine, aspartic acid, and a mixture of glycine and aspartic acid, all 0.03 M, will be used. If desired, a fourth solution containing either or both amino acids can be used as an unknown.

Cut strips of Whatman No. 1 filter paper to the dimensions 13.5 \times 1.8 \times 1.0 cm. Pierce a small hole in each strip, in the center and about 4 mm. from the broad end. This will he used for hanging the strips during the drying process. To minimize contamination, avoid touching the strips with the fingers.

For each solution under examination, draw a portion into a capillary pipet having a very fine tip,4 and apply the tip of the pipet quickly and lightly to a dilter-paper strip at the center and about 6 mm. from the narrow end. By this procedure a volume of approximately 0.2 µl. of solution is deposited on the paper to form an area which should not be in excess of 1.5 mm. In diameter. Circle the wet area with a light pencil line and allow to dry. Prepare a separate strip for each solution under examination.

Place approximately 0.5 ml. of water-saturated phenol! in the bottom of a 6-inch test tube. Using one tube for each strlp, insert the narrow end of the treated paper strlp into the phenol solution in such a way that the strlp does not touch the walls of the tube except at the top. Stopper the tube with a soft cork. Allow to stand undisturbed at room temperature for 2 to 3 hours or until the solvent has ascended by capillarity to within about 5 mm. of the hole at the top of the strlp. Remove the strlp, place a bent wire or paper ellp in the hole, and suspend in a drying oven at 110° for about 3 minutes. Remover, and spray the entire strlp lightly with a solution of ninhydrin contained in an atomizer. Replace in the oven and dry for about 4 minutes.

With a pencil, mark the edge of the solvent front on the strip, and encircle any colored spots present. Mark the approximate center of each spot. Measure the distance from the original spot where the solution was applied to the edge of the solvent front and to the center of each colored area. Calculate the Ry value for each colored spot, where Ry is the ratio of the distance traveled by the material in the spot to the distance traveled by the solvent. How do your values compare with those given by Rockland and Dunn (glycine, 0.49; aspartic acid, 0.25)? From the volume of solution used, and the moiar concentration of smino acid, calculate the actual amount of amino acid separated and identified by this procedure.

Do not overlap the edges, but allow the staples to serve as links Stand the cylinder upright, with the applied spot at the bottom, in a 10 cm petri dish filled to a depth of about 0 5 cm with a suitable solvent (see below). Cover the assembly with a bell jar or battery jar, preferably with a tightly fitting glass plate as a base. Allow to stand until the solvent has risen to within a few centimeters of the top of the cylinder Remove from the petri dish, dry quickly in an oven, and unroll Remove the staples, turn the paper so that the circled spot is in the lower right corner, and make a new cylinder as before Place upright in a petri dish containing a second solvent, cover with a bell jar, and let stand as before. When the solvent has risen to within a few centimeters of the top of the paper, remove from the petri dish and dry quickly in an oven. Unroll, spray the dried paper uniformly with ninhydrin solution, and dry as described under Exp. I. Circle the colored areas with a pencil line to mark the spots, since the colors will fade in a few days.

Try this experiment with solutions of various amino acids and with various solvents. Suitable solvents! include (1) water-saturated phenol (Exp. 1), (2) a mixture of 70 parts n-propanol and 30 parts water, (3) a mixture of equal parts of 2,4,6 collidine, and 2,4 lutidine to which one third volume of water is added, (4) a mixture of test butanol, water and 85% formic acid in the proportions of 69 5 29 5 1 0, (5) water saturated phenol made alkaline with 1% concentrated NH₄OH 110w do your results compare with those shown in the frontisplece?

- 3 Detection of Glutomote in Canned Foods An Interesting use of the procedure described in Exp 1 has been described by Patton and Foreman's for the detection of added sodium glutamate in processed foods Sodium glutamate is an approved flavoring agent which may or may not be added to canned foods such as chicken soup Try the procedure of Exp 1, using fluid from commercial canned chicken soup or meat sauce. As a control, run an 003 M solution of sodium glutamate. As a second control, glutamate may be added directly to the solution under test. Do your results indicate the presence of glutamate in the food tested?
- 4 Column Chromotograph; Experiments Illustrating the use of column chromatograph; will be found in Chapters 7 and 10

EXPERIMENTS ON CHROMATOGRAPHY

I One dimensional Paper Chromatography * For demonstration purposes solutions of amino acids will be chromatographed in this experiment. The amino acids will be located on the filter paper by reaction with ninhydrin to give a purple color. Solutions of glycine, aspartic acid, and a mixture of glycine and aspartic acid, all 0.03 M, will be used II desired, a fourth solution containing either or both a mino acids can be used as an unknown.

Cut strips of Whatman No 1 filter paper to the dimensions 13 5 × 18 × 10 cm Pierce a small hole in each strip, in the center and about 4 mm from the broad end This will be used for hanging the strips during the drying process To minimize contamination, avoid touching the strips with the fingers

For each solution under examination, draw a portion into a capillary pipet having a very fine tip 4 and apply the tip of the pipet quickly and lightly to a dilter paper strip at the center and about 6 mm from the narrow end By this procedure a volume of approximately 0 2 al of solution is deposited on the paper to form an area which should not be in excess of 1 5 mm in diam eter Circle the wet area with a light pencil June and allow to dry Prepare a separate strip for each solution under examination

Place approximately 9 5 ml of water saturated phenol* In the bottom of a 6 inch test tube. Using one tube for each strip, insert the narrow end of the treated paper strip into the phenoisolution in such a way that the strip does not touch the walls of the tube except at the top. Stopper the tube with a soit cork. Allow to stand undisturbed at room temperature for 2 to 3 hours or until the solvent has ascended by capillarity to within about 5 mm of the hole at the top of the strip. Remove the strip, place a bent wire or paper clip in the hole, and suspend in a drying oven at 110° for about 3 minutes. Remove, and spray the entire strip lightly with a solution of ninhydrin contained in an atomizer 8 Replace in the oven and dry for about 4 minutes.

With a pencil mark the edge of the solvent front on the strip, and encircle any colored spots present Mark the approximate center of each spot Measure the distance from the original spot where the solution was applied to the edge of the solvent front and to the center of each colored area. Calculate the Ry value for each colored spot, where Ry is the ratho of the distance traveled by the material in the spot to the distance traveled by the solvent How do your values compare with those given by Rockland and Dunn (glycine, 0.49 aspartic acid, 0.25) From the volume of solution used, and the molar concentration of similar solvents of similar solvents and the solvents are acid and identified by this procedure.

Do not overlap the edges, but allow the staples to serve as links. Stand the cylinder uprlight, with the applied spot at the bottom, in a 10-cm, petri dish filled to a depth of about 0.5 cm. with a sultable solvent (see below). Cover the assembly with a bell jar or battery jar, preferably with a tightly fitting glass plate as a base. Allow to stand until the solvent has risen to within a few centimeters of the top of the cylinder. Remove from the petrl dish, dry quickly in an oven, and unroll. Remove the staples, turn the paper so that the circled spot is in the lower right corner, and make a new cylinder as before. Place upright in a petri dish containing a second solvent, cover with a bell jar, and let stand as before. When the solvent has risen to within a few centimeters of the top of the paper, remove from the petri dish and dry quickly in an oven. Unroll, spray the dried paper uniformly with ninhy drin solution, and dry as described under Exp. 1. Circle the colored areas with a pencil line to mark the spots, since the colors will fade in a few days.

Try this experiment with solutions of various amlno acids and with various solvents. Suitable solvents include (1) water-saturated phenol (Exp. 1); (2) a mixture of 70 parts 11-propanol and 30 parts water; (3) a mixture of equal parts of 2,4,6-collidine, and 2,4-iutidine to which one-third volume of water is added; (4) a mixture of tert.-butanol, water and 85% formic acid in the proportions of 69,5:29.5:1.0; (5) water-saturated phenol made alkaline with 1% concentrated NH₄OII. How do your results compare with those shown in the frontispiece?

- 3. Detection of Glutamate in Conned Foods. An interesting use of the procedure described in Exp. 1 has been described by Patton and Foremanis for the detection of added sodium glutamate in processed foods. Sodium glutamate is an approved flavoring agent which may or may not be added to canned foods such as chicken soup. Try the procedure of Exp. 1, using fluid from commercial canned chicken soup or meat sauce. As a control, run an 0.03 M solution of sodium glutamate. As a second control, glutamate may he added directly to the solution under test. Do your results indicate the presence of glutamate in the food tested?
- 4. Column Chromatography. Experiments illustrating the use of column chromatography will be found in Chapters 7 and 10.

OSMOTIC PRESSURE

Membranes which permit the passage of molecules of the solvent but not of the molecules or ions of a substance in true solution are known as semipermeable membranes. One type of semipermeable membrane is made by depositing copper ferrocyanide in the walls of a porous porcelain cup If such a cup is filled with a solution of sugar in water and is then immersed in water, the sugar molecules cannot pass through the membrane. The water molecules, however, permeate readily and the level of the solution in the cup will rise. The pressure that must be exerted upon the solution within the cup to prevent any increase in volume is a measure of the osnotic pressure of the solution. The magnitude of osnotic pressure is indicated by the fact that a 10 per cent solution of cane sugar at 25° C, has an osmotic pressure of 7.6 atmospheres.

Whatever may be the fundamental cause of osmotic pressure there is no

^{*} Underwood and Bockland Food Res. 18, 17 (1953).

¹⁵ Patton and Foreman, Food Tech., 4, 83 (1930).

inward attraction is that the number of molecules in the surface tends to be reduced to a minimum and the surface contracts until its area is the smallest possible for a given volume of liquid. This explains the tendency of small drops of liquid to assume a spherical form. This spontaneous contraction of a liquid surface indicates that there is a certain free energy in the surface, and that in order to extend this surface by hringing molecules into the surface a certain amount of work must be done in overcoming the attraction of the other molecules in the interior. It is customary, in considerations involving surface forces, to substitute for the free surface energy the concept of a surface tension acting in all directions parallel to the surface. Thus, if the surface tension of a liquid is γ dynes per em, the work done in increasing the surface area by 1 sq em will be γ ergs, and the free surface energy of the surface will be γ ergs per sq cm.

When two immiscible liquids are in contact with each other, the dividing surface or interface also contracts spontaneously because each type of molecule in the surface is attracted by its own kind of molecule in the interior. Thus the interface also possesses a free surface energy, or a surface tension known as the interfacial tension. When the interfacial tension between two liquids is zero, the two liquids are completely miscible.

An intimate mixture of two immiscible liquids is known as an emulsion, one liquid heigh dispersed in the form of droplets in a continuous or external phase of the other liquid. Dimilisfication is facilitated by the presence of substances called emulsifying agents, which minimize the tendency of the droplets to coalesce. Among the most effective of such agents are substances which lower surface tension, such as soaps, proteins, bile salts, etc. The action of hile salts in lowering surface tension is an important aid in the emulsification and digestion of fats in the intestinal tract

Emulsions may be of either the oil-in water or the water in-oil type, depending upon which liquid is the continuous phase Sodium soaps, which are more soluble in water than in oil promote the formation of oil-in-water emulsions, calcium or magnesium soaps, which are oil-soluble promote the formation of water-in-oil emulsions. This has been offered as an explanation for certain antagonistic effects of Na+ and Ca++ ions on the physical state of protoplasm.

2. Determination of Relative Surface Tension by Drop Method. The surface of a liquid tends to contract so as to produce a minimum area. When flowing through a small opening it therefore tends to form drops which fall away when gravity overcomes the surface tension. The number of drops formed from a given weight or volume of fluid varies, therefore, with the surface tension.

Procedure. Set up a Traube stalagmometer as in Fig. 3 so that the whole may be immersed in a constant-temperature bath. For approximate com-

parative results an ordinary 1- or 2-ml. pipet elamped in a vertical position may be used, regulating the outflow by the use of the finger or n piece of rubber tubing with attached screw clamp so that discrete drops are formed at a uniform rate, $Fill\ lt$ with water above mark A. When the level of the water falls to this mark, start counting the number of drops that fall until mark B is reached, By means of the scale divisions above and below A and B (knowing how many divisions correspond to a drop), fractions of a drop may be estimated. If γ represents the relative surface tension of the liquid tested, $N_{\rm BF}$ the number of drops of water, N the number of drops of the unknown, and S its specific gravity, $\gamma = \frac{N_{\rm BF} \cdot S}{N_{\rm BF} \cdot S}$

Determine the surface tensions of water, of a very dilute soap solution, and of olive oil, cleaning the instrument thoroughly between tests, and rinsing several times with the solution tested before taking readings. Olive oil has a lower surface tension than water. Soaps markedly lower the surface tension of aqueous solutions.

3. Determination of Surface Tension by Ring Method of Du Nolly. The drop method for determining surface tension described in Exp. 2 is a static method

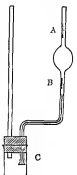


Fig 3 Stalaguometer

So also is the measurement of the rise of fluid in a capillary tube. The osciliating-jet method is a dynamic method in that new surfaces are constantly being formed. All methods give practically concordant results in the case of pure liquids, but not in the case of solutions. "Du Nouy has suggested a static method based on the measurement of the force required to overcome the adherence of a platinum ring to the surface of the liquid in question. The apparatus consists of a platinum ring hanging from a lever arm which can be raised by applying torsion to the wire to which it is fixed. The torsion is measured on a scale which can be calibrated directly in dynes per cm., the absolute unit of surface tension. Determinations of surface tension by this method may be accurately made on 1-ml. portions of fluid in 20 to 30 seconds. The method is particularly useful in physiological studies. "The surface tension."

que-tion but that it appears to be, for sub-tances in solution, analogous to the pressure of gas-cous molecules. Thus 1 gram-molecular weight of an ideal gas at 0°C and in a volume of 1 liter everts a pressure of 22.4 atmospheres; 1 gram-molecular weight of an ideal nonelectrolyte in solution under the same conditions has an osmotic pressure of 22.4 atmospheres. Furthermore under ideal conditions osmotic pressure is, like gas pressure, directly proportional to the als-olute temperature and to concentration, and is independent of the chemical nature of the dissolved material. Thus equimolecular conventrations of all nonelectrolytes have the same osmotic pressure, or are isosmotic. It is therefore possible to determine molecular weight by the measurement of osmotic pressure, and this procedure has found considerable application in biology.

The osmotic pressure of electrolytes is considerably higher than that

and ostation presents at electronics is considerany higher than that of equimolar solutions of nonelectrolytes, since an ion is theoretically as effective o-motically as a molecule. For example, a solution of NaCl containing Na* and Cl- ions should have twice the o-motic pressure of an equimolar solution of nonelectrolytes such as glucoce or urea. By actual measurement it is usually found that the o-motic pressure of electrolytes is somewhat less than the expected value, presumably because of differences between the o-motic effectiveness (actually) of the ions and their concentration. The relation between ion concentration and activity must be considered in any calculation of o-motic pressure for solutions containing electroly terms.

A colloidal particle is as effective comotically as a molecule or ion. Colloidal solutions therefore have very low comotic pressures compared to equal weights of substances in true solution because of the large size and relatively small number of particles present. Colloidal osmotic pressures, although small, are nevertheless a very important in biology, since colloids do not ordinarily diffuse through membranes and only substances which are impermeable to a membrane ordinarily can influence the osmotic flow of fluid across the membrane. For example, the osmotic pressure of the blood colloids is only about 0.5 of 1 per cent of the total osmotic pressure of blood, yet the blood colloids become the determining factor in the flow of fluid by osmosis across those physiological membranes which are freely

follow; but for practical reasons associated with the enormous pressures concerned and the difficulty in obtaining a truly semipermeable membrane, other methods are usually employed for all except colloidal solutions. These methods make use of those other physical properties of solutions which, like osmotic pressure, depend upon the number of dissolved particles (e.g., vapor pressure, rise in boiling point, or depression of freezing point of the solvent). An aqueous solution with an osmotic pressure of 22.4 atmospheres, for example, has a freezing point 1.86° C. below that of pure water, with other values in direct proportion. The freezing-point method is by far the most widely used in investigations on osmotic pressure in biological fluids.

EXPERIMENTS ON OSMOTIC PRESSURE

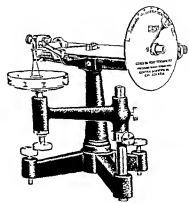
- 1. Demonstration of Osmotic Pressure of a True Solution (Cone Sugor). Prepare a dialyzing bag similar to that described on p. 10. The the mouth of the bag to a one-hole rubber stopper. With a pipet fill the bag with a 10 per cent solution of cane sugar. Insert a glass tube (about 30 cm, long and of 1 to 2 mm, bore) through the hole in the stopper until the end dips below the surface of the fluid in the bag. Blow gently through the tube to create a slight positive pressure in the bag. Support the bag in a large beaker of water with the glass tube in an upright position and the level of fluid in the bag and beaker approximately the same. Observe the rise of fluid in the glass tube. Ultimately the sugar will diffuse through the membrane and the solution level in the tube will fail, but meanwhile the existence of an osmotic pressure will be demonstrated. Copper ferrocyanide membranes deposited in porcelain cups are truly semipermeable, and are necessary for quantitative determination of the osmotic pressure of true solutions by this method.
 - 2. Demonstration of Osmatic Pressure of Colloidal Solutions. The bag employed in Exp. 1, or a similar one, may be used and the determination is made in the same way. The highest point of the column is generally reached in five or six hours. Memhranes of this type are not permeable to proteins, so a true measure of their osmatic pressure may be obtained. The pressures attained, however, are very much lower. A 1 per cent solution of gelatin may give a rise of 25 to 50 mm.
 - 3. Osmotic Pressure of Red Blood Cells. Add a drop or two of blood to 5 ml, of each of the following solutions: 0.3, 0.9, and 5.0 per cent sodium chloride. Examine each suspension under the microscope. Explain the findings. Caiculate the molar concentration and approximate osmotle pressure of each of the three solutions. What is the osmotle pressure of the red-cell contents?

Repeat, using the following solutions: 0.16 M sodium chloride, 0.32 M urea, and 0.32 M urea containing 0.16 M sodium chloride. Observe as above. Calculate the osmotle pressure of these solutions. Which are isosmotic with the red cell? Which are isosmotic? Explain.

SURFACE TENSION

The molecules in the interior of a liquid are surrounded by other molecules on every side and are thus subject to uniform attractions in every direction. On the other hand the molecules in the surface are attracted inward and by other molecules in the surface, but there is little attraction outward because there are so few molecules outside. The result of this

sion of a resting surface of serum diminishes on standing (time drop) due to the gradual accumulation of surface active substances (see p. 22). This phe nomenon has been investigated by Du Nouy. "Using this apparatus, determine the surface tension of the liquids tested in Exp. 2 and compare the results obtained by the two methods (see Fig. 4). This apparatus has been modified to permit the measurement of interfacial tension between two figuids."



and "oil" phases of protoplasm has attracted considerable attention. It is not surprising to find that substances which are predominantly lipide-soluble are concentrated in the various fats and oils of living tissues. The presence of fat-soluble vitamins in various fish-liver oils is an example, as is also the reported concentration of the insecticide DDT in the fat of milk obtained from cows exposed to DDT-contaminated areas. The action of drugs has been studied in relation to water-lipide distribution. According to one theory, the relative lipide-solubility of various drugs is of importance in determining penetrability of such drugs into, and action upon, such tissues as the brain and nerves, which are relatively rich in lipide material. In connection with such theories however it must be remembered that the physical nature of the so-called oil phase of protoplasm has not been too clearly defined. There are relatively few tissues other than adipose and similar tissue where discrete fat droplets are found within the cell. The lipide material of brain and nerve tissue is not fat or oil but consists of phospholipides and similar compounds (see p. 290) which are dispersed within the primarily aqueous phase of the cell in a way which is not entirely clear, but is known to involve intimate association with the proteins and other water-soluble constituents of the cell. The extent to which the distribution of a drug or other substance between water and peanut or olive oil, as determined in the laboratory, is applicable to the complex environment of the living cell remains to be more clearly defined.

The distribution of dissolved material between immiscible solvents has many practical applications. Widespread use is made of such nonaqueous solvents as ether, chloroform, and benzene for the extraction from biological material of lipides, fat-soluble vitamins, various organic acids, and other similar substances. An important advance in the use of solvent distribution for the isolation and characterization of substances of biological importance has been made by Craig's in devising his procedure for countercurrent distribution. In this procedure the material under examination is subjected to a unique and systematic series of extractions by an immiscible solvent in a sequence of extraction vessels or tubes in such a way that various substances become separated and distributed among the

of impurities Countercurrent distribution has found wide application in the isolation and characterization of biologically active substances, comparison of natural and synthetic compounds, and establishment of the purity of drugs and similar substances An experiment illustrating the principles involved is described below.

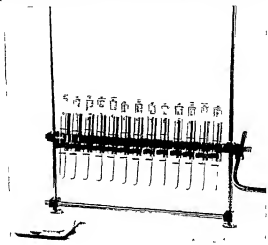
EXPERIMENTS ON DISTRIBUTION OF DISSOLVED SUBSTANCES BETWEEN LIQUID PHASES

Countercurrent Distribution. 1. This experiment is designed to illustrate the principles of Crafg's countercurrent-distribution procedure for the isolation and identification of substances in solution. A study will be made of the distribution of propionic acid between 22 M phosphate buffer at pH 52 (aqueous phase) and isopropyl ether.

Set up nine 25.m.l graduated cylinders with tightly fitting glass stoppers, and number them from 0 to 8. Place 15 ml. of water-saturated isopropyl ether? Into each tube Into Tube 9 place 0.2 ml. of propionic acid. Now add 7.5 ml of phosphate buffer to Tube 0, stopper, and invert 50 times. Allow the layers to separate. Remove the lower layer as completely as possibles and transfer it to Tube 1, Add 7.5 ml. of fresh lower phase to Tube 0. Stopper, equilibrate by inverting hoth tubes 50 times as before, and allow the phases to separate. Transfer the lower phase of Tube 1 to Tube 2, the lower phase of Tube 0. Again Invert all tubes 50 times. Continue the transfer of lower phase to Tube 0. Again Invert all tubes 50 times. Continue the transfer of lower phase to Tube 0, and the equilibration, until all nine tubes have equal volumes of both phases. The distribution is now completed within the limits of the number of tubes used. During this scoreiment, the volumes of the two phases should be noted

occasionally, particularly that of the leading lower phase, and any deficit made up by the addition of fresh lower or upper phase as required, so that the ratio of phases is maintained at 2:1 in each tube To facilitate equilibration between transfers, it is convenient to clamp the cylinders or tubes to a horizontal rod which can be turned with a crank. Such a derice is illustrated in Fig. 5 in place of the glass cylinders, test tubes or separatory funnels may be used.

drops of 1 per cent alcoholic phenolphthalein solution to each flask, and titrate with 0.01 N NaOH solution to the first permanent pink color. As a control, titrate 2 ml. of fresh upper phase in the same way. Subtract the buret reading for the control from that for each tube. Plot milliliters of alkali required for each tube against tube number, on cross-section paper, and draw



upper-phase volume to lower-phase volume, i.e., $r\approx 2.0$ How does your value compare with the value of 0.50 estimated by Sato et al.?

Another method for calculating the value of K is based on the fact that the ratio of the concentrations of substance analyzed for in any two tubes is related to K as follows:

 $K = \frac{1}{r} \times \frac{T_N}{T_{N-1}}$

where T_N and T_{N-1} refer to the amounts of material (or aliquots thereof, since ratios are concerned) found in tube X and in the tube immediately preceding it in the countercurrent distribution F is a factor, differing for each tube, if with the following values in this experiment: N=2, F=24, X=3, F=55, X=4, F=54, X=5, F=55; X=6, F=55; X=6, F=56; X=6; X

$$K = \frac{1}{r} \times F \times \frac{T_{N-1}}{T_{n-1}}$$

Using this equation, calculate K when N equals 2, 3, 4, 5, and 6. Average your results llow do they compare with the value obtained above?

2. Repeat this experiment using (a) acetic acid (K=0.09) and (b) butyric acid (K=2.04). How do the curves and calculated results compare with those for propionic acid?

HYDROGEN-ION CONCENTRATION

Acidity. Acids may be defined as compounds which yield positively charged hydrogenions²⁴ in solution; bases, as compounds which yield negatively charged hydroxyl ions in solution Strong ands, such as HCl, are completely ionized at all concentrations; they therefore have a hydroxylionized at all concentration and acid present. Weak acids, such as accetic acid, evist in solution largely in the molecular or undissociated form, and have a hydroxylionized concentration and evidinary conditions which is small relative to the total acid concentration. A similar distinction is made between strong bases and weak bases. Nost neutral salts are considered to be completely ionized in solution.

When acids react with bases a double decomposition occurs which results in the formation of a salt and nater, as in the following equation.

$$HA \rightleftharpoons H^+ + A^-$$

Such reactions, known as neutralization reactions, go to completion because the water formed during the reaction is itself so feebly ionized that its formation leads to the removal of practically all of the hydrogen and hydroxyl ions from solution. The amount of base required to neutralize a definite volume of acid (i.e., the titratable acidity) depends entirely upon the concentration of the acid and is independent of its degree of dissociation. This fact becomes evident upon study of the above equation. As the base is added some of the hydrogen ions in the solution combine with the added hydroxyl ions to form water, and are thus removed from solution. This removal of hydrogen ions disturbs the equilibrium between the undissociated molecules of acid and its ions, and more of the acid dissociates in an attempt to restore this equilibrium. If sufficient base is added this process will continue until all of the acid has been dissociated and neutralized. This will be the case irrespective of whether the acid was originally highly ionized or feebly ionized—i.e., whether it was a strong acid or a weak acid. On the other hand, the hydrogen-ion concentration of an acid-i.e., the actual amount of free hydrogen ions present in the solution at a particular time-depends not only upon the concentration of the acid but also, for weak acids, upon the degree of dissociation. Thus the hydrogen-ion concentration of normal hydrochloric acid, which is 100 per cent ionized, is approximately 100 times as great as the hydrogen-ion concentration of normal acetic acid, in which less than 1 per cent of the molecules are dissociated. As indicated above, however, equal volumes of normal hydrochloric and normal acetic acids will be neutralized by exactly the same amounts of base. In other words the titratable acidities of these acids, or the amounts of hydrogen ions which they are capable of yielding on complete dissociation, are the same. The preceding discussion of the hydrogen-ion concentrations and titratable acidities of weak and strong acids applies just as well to the hydroxyl-ion concentrations and the titratable alkalinities of weak and strong bases.

Chemical Equilibrium: Ionization Constants. According to the law of mass action a reversible reaction of the type

$$A + B \rightleftharpoons C + D$$

proceeds from left to right at a velocity (v_1) which is proportional to the product of the concentrations of A and B—i.e., $v_1 = k_1[A][B]$. Similarly, the opposite reaction takes place at a rate (v_2) proportional to the product of the concentrations of C and D—i.e., $v_2 = k_2[C][D]$. At equilibrium the velocities in opposing directions are equal, and hence $k_1[A][B] = k_2[C][D]$

or $\frac{[C][D]}{[A][B]} = \frac{k_1}{k_1} = K$. This expression of the law of mass action states that for a reversible reaction the ratio of the product of the concentrations of the reacting substances on one side of the equation to that of the product of the concentrations of the substances on the other side is constant. That is,

$$\frac{[C][D]}{[A][B]} = E$$

Applying this to weak acids, HA, which ionize according to the equation, $HA \rightleftharpoons H^+ + A^-$, we find that at equilibrium

$$\frac{[H^+][A^-]}{[HA]} = K_A$$

In this case the equilibrium constant, since it measures the equilibrium between the undiscocated molecules of icid and its ions, is known as the dissociation constant, K_{κ} . For weak bases, similarly, the dissociation constant of the base BOH is given by

$$\frac{[B^+][OH^-]}{[BOH]} = K_B$$

From the above equations it is readily seen that the larger the numerical value of the dissociation constants, the more completely the acid or base is dissociated—i.e., the stronger the acid or hase

An acid solution is one that contains an excess of hydrogen ions. An alkaline solution is one that contains an excess of hydroxyl ions. A neutral solution is one that contains hydrogen and hydroxyl ions in equal concentrations. Pure water is a neutral solution. It is dissociated to an extremely small extent according to the equation.

Applying the law of mass action to this dissociation we get

$$\frac{[II^+][OII^-]}{[II_2O]} = K$$

Since the concentrations of \mathbb{H}^* and \mathbb{OH}^- in pure water or in dilute aqueous solutions are so small compared with the concentration of un dissociated water molecules, the concentration of the latter, $[H_2O]$, may be considered as constant, and the equation then becomes $[\mathbb{H}^+] \times [\mathbb{OH}^+] = K [\mathbb{H}O] = K_*$ in other words, the product of the concentrations of the \mathbb{H} and \mathbb{OH} income is constant. The value of K_m at 25° \mathbb{C} has been found to be 1×10^{-11} . That is, $[\mathbb{H}^+] \times [\mathbb{OH}^+] = 1 \times 10^{-11}$ in pure water, for every hydrogen ion set free a hydroxyl ion must also be liberated so that the concentrations of the two ions remain equal—that is, $[\mathbb{H}^+] = [\mathbb{OH}^+] = [\mathbb{OH}^+] = [\mathbb{OH}^+] = [\mathbb{OH}^+] = 1 \times 10^{-11}$. Then $[\mathbb{H}^+] = [\mathbb{OH}^+] = 1 \times 10^{-11}$. Thus pure water or a neutral solution contains approximately $1 \times 10^-$ moless** of \mathbb{H} or \mathbb{OH} ions per liter and is a 1/10 00 000 normal solution of \mathbb{H} or \mathbb{OH} ions per liter and is a 1/10 00 000 normal solution of \mathbb{H} or \mathbb{OH} in \mathbb{OH} .

Hydrogen ion concentrations are now generally expressed, for practical as well as theoretical reasons as their logarithms with the sign reversed and indicated by the term pH Γhus^{22} pH = $-\log[H^+]$, or [H⁺] = 10^{-pR}

The logarithm of 1×10^{-7} is -7.0 and the pH is 7.0. To take another example, the hydrogen-ion concentration of 0.1 N HCl is 8.3×10^{-2} . The logarithm of the product of two numbers is equal to the sum of their respective logarithms. In this ease, then, $\log (8.3\times 10^{-2}) = \log 8.3 + \log 10^{-2}$ (consulting table of logarithms²⁴) = 0.92 + (-2) = -1.03. The pH, being the negative log of the hydrogen-ion concentration, equals -(-1.08) or 1.03. To convert pH values to hydrogen-ion concentrations the procedure is reversed—e.g., in the case of pH 1.03, the $[\mathrm{H}^+] = 10^{-10}$ s $= 10^{-24} \, \mathrm{y}^{2} = 10^{-2} \times 10^{9} \, \mathrm{z}^{2}$. The logarithm of a number is the power to which 10 must be raised to give the number, hence the value of $10^{0.92}$ is obtained by looking up the number whose logarithm is 0.92 (i.e., antilog 0.92). This is found to be 8.3. pH 1.08 is therefore equal to a $[\mathrm{H}^+]$ of 8.3×10^{-2} .

Since the product of the H and OH ion concentrations is constant at 1×10^{-4} , when the [H+] increases from 1×10^{-7} (pH 7.0) to 1×10^{-4} (pH 4.0) the [OH-] decreases to 1×10^{-10} (pOH 10.0). The sum pH + pOH always equals 14. According to this nomenclature an increase in pH indicates a decrease in hydrogen-ion concentration or true acidity.

Since the hydrogen-ion concentration varies in a definite reciprocal manner with hydroxyl-ion concentration, the pH scale is universally used to express degrees of alkalinity as well as of acidity, as indicated in the second column below.

Normality of [H+]	pН	}	HOq	Normality of [OH-]
N/10 N/1,000 N/1,000,000	1 3 6	Aendity	13 11 8	
N/10,000,000	7	Neutrality	7.7	N/10,000,000
	8 11 13	Alka- limty	6 3 1	N/1,000,000 N/1,000 N/10

Buffer Action and Buffers. By buffer action is meant the ability of solution to resist marked change in pH on the addition or loss of acid these Substances whose presence in the solution are responsible for uffer action are known as buffers. Though strong acids or bases may ometimes act as buffers, under ordinary conditions the most common asis for buffer action lies in the presence within the solution of a weak cid (or weak base) together with its salt. Such solutions have characeristic properties with respect to hydrogen-ion concentration which are of value not only in the laboratory but also in the living organism, and an inderstanding of these properties is of fundamental importance.

If we take for example a solution containing a weak acid, HA, and its salt, BA, the hydrogen ions in the solution can come only from the lissociation of the acid molecules The dissociation coulibrium of the

¹⁴ See Appendix.

acid is given by the equation $\frac{[H^+][A^-]}{[HA]} = K_A$, as described previously.

From this it follows that the hydrogen-ion concentration [H⁺] $\approx K_{A} \frac{[\text{HA}]}{[\text{A}^{-}]}$

For a weak acid, which is relatively little ionized, the difference between the concentration of undissociated acid molecules [HA] and the total acid concentration [Acid] is small. This difference becomes even less in the presence of a salt of the acid, since the salt ions A-, by a mass action effect, repress still further the dissociation of acid molecules. Thus in a mixture of salt and acid we can substitute the total acid concentration [Acid] for the quantity [HA], with very little error. Similarly the concentration of acid ions A- derived from the dissociation of the acid molecules is very small compared to that furnished by the salt BA which is completely dissociated into B+ and A- ions, so that we may consider the value of [A-] to be determined primarily by the salt concentration Although the salt is completely dissociated, because of the existence of interionic forces the effective concentration of salt ions, [A-], is some what less than the actual salt concentration. The ratio between these two quantities is, however, sensibly constant over the range of concentration that ordinarily comes into question, so that we may say [A-] equals ESalt]. Substituting ESalt] for [A-] in the above equation, and combining constants, we obtain the expression

$$[H^+] = K'_A \frac{[Acid]}{[Salt]}$$

This equation tells us that the hydrogen-son concentration of a solution containing a weak acid and its salt is determined by the value of K', for the acid and by the ratio of acid concentration to salt concentration in the solution. While usable in the form given, it is more convenient to convert it into terms of pII rather than of [II+] If the logarithm of both sides of the equation is taken, and the signs reversed, we obtain:

$$\sim \log \{H^+\} = -\log K'_A - \log \frac{[Acid]}{[Salt]}$$

Now $-\log [H^*]$ has already been defined as equal to pH. In an analogous way and for similar reasons we can define a term pK'_A as equal to $-\log K'_A$, and we can replace the negative expression $-\log \frac{[Acid]}{[Scalt]}$ by

its equivalent, the positive expression $\log \frac{[Salt]}{[Acid]}$. Substituting the various terms in the above equation, we obtain:

particular acid, and by the logarithm of the ratio of salt concentration to acid concentration in the solution For example, the value of K_A for acid is approximately 1.8×10^{-5} , or $10^{-4.7}$ The p K_A for acid is therefore 4.7, and the pH of an acetic-acid-sodium-acetate buffer is given by the equation

 $pH = 4.7 + \log \frac{[Sodium\ acetate]}{[Acetic\ acid]}$

When [Sodium acctate] = [Acetic acid] the pH is 47, since the ratio of salt to acid is 1, and $\log 1 = 0$ By varying the proportion of salt to acid in the mixture the pH will vary over a range of approximately 15 pH units below and above the pK'_A value—1e from pH 32 to pH 62 Outside of this range the equation is not applicable for this particular system. To cover other pH ranges other weak acids with suitable pK'_A values are selected, so that by this means it is possible to prepare huller solutions of almost any desired pH. This procedure is illustrated in the section on the pieparation of pH standards which follows

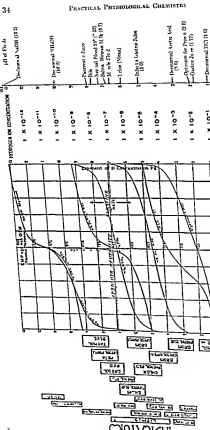
In addition to their value in the preparation of solutions of known pH, buffer solutions possess another property of fundamental importance in the high granism, namely the ability to resist marked changes in pH on the addition of small amounts of strong acids or bases. It has been pointed out by Van Slyke that if a small amount of strong acid such as HCl is added to blood which contains several buffers the resulting hange in pH in the direction of increased readity may be only 1/1,000 of hat which would have occurred if the blood were unbuffered. The action of a buffer solution in imminizing the effect on pH of added acid or base any be visualized as follows.

When hase is added to the mixture the excess hydroxil ions are renoved by the hydrogen ions coming from the neid, which combine with hem to form water. Upon the addition of acid the excess hydrogen ions from the added acid are removed by the salt ions, which combine with them to form more molecules of the relatively un ionized weak acid. The reactions myolved are as follows.

Buffer Acid $HA \rightleftharpoons H^+ + A^-$ on addition of base, $OH^- + H^+ \rightarrow H \cdot O$ Mixture Salt $BA \rightleftharpoons B^+ + A^-$ on addition of acid $H^+ + A^- \rightarrow HA$

Thus the addition to a buffer solution of small amounts of base results merely in the production of more salt ions at the expense of an equivalent amount of the weak acid present the addition of acid similarly results in the production of more weak acid at the expense of the salt The pH of the solution must inevitably change, but this change in accordance with the demands of the Henderson-Hasselbalch equition is due to a change in the value of the logarithm of the ratio of salt concentration to acid concentration which innerically is quite small in comparison to the concentration of II+ or OH- ions added to the solution

It is obvious from what has just been said that the more concentrated a buffer is the smaller will be the change in pH on the addition of a given amount of strong and or base—je the greater will be the buffer power of the solution. Furthermore, if for example and is added to a buffer solu-



Parts of the less acid constituent

1 × 10-1

This chut is, in slightly modified form, one prepared by Walpole from a large amount of data accumulated by Serenson, Walpole hinwelf, Alchenia, and a number of other authors See Najpole Biochem J., 8, 624 (1911) Indicators marked * are used in intrational but not for plf determinations.

tion in an amount greater than that equivalent to the buffer salt present, the buffer capacity of the system will be exceeded and its ability to function as a buffer will disappear, since it will no longer consist of a mixture of buffer salt and acid. Another property of buffer solutions which is also evident from inspection of the Henderson-Hasselbalch equation, is that they may be diluted considerably without appreciable change in pH, since the pH depends upon a concentration ratio rather than upon concentration itself.

The above discussion has been based entirely upon a consideration of buffers containing weak acids and their salts. An entirely analogous derivation can be made for solutions containing weak bases and their salts.

Preparation of Standard Buffers of Known pH. Fig 6 indicates how standard solutions of definite hydrogen ion concentration may be made up from a series of stock solutions, mixed in definite proportions The stock solutions indicated on the chart were suggested by Sørensen and are as follows 0 10 N HCl 0 10 N NaOH, 7 505 g gly one plus 5 85 g NaCl per liter, 11 876 g Na-HPO, 2H₂O per liter, 25 9 078 g kH-PO, per liter 21 008 g eithe acid in 1 liter of 0 20 N NaOH, 12 404 g boric acid in 1 liter of 0 10 N NaOH The other solutions are 0 20 N sodium acetate and 0 20 N acetic acid. Solutions of known hydrogen ion concentration are prepared from these by mixing in the proportions indicated in Fig. 6, the abscissas representing parts of the relatively more alkaline constituent. Thus a mixture of seven volumes of the sodium acetate stock solution with three volumes of the stock acetic acid solution gives a mixture with a hydrogen ion concentration of 1×10^{-5} (pH 5 0) The mixtures are most satisfactory through the ranges where the hy drogen ion concentrations change most gradually—that is through the flatter portions of the curves. The phosphate mixtures covering the range 53 to 80 are especially useful in biological work. The standard huffer solutions of Clark and Lubs, described below, are arranged to differ by even increments in pH instead of in component solutions, and are quite

0.2 M KCl add the indicated number of ml of 0.2 N HCl and dilute to 200 ml Indicator thymol blue

Hq	HCI	pH	IICI	pH	11C1	pH	1101
1 2 1 4	64 5 41 5	1 6 1 8	26 3 16 6	2 0	10 6	2 2	6 7

Group 2 To 50 mt of 0.2 M acid potassium phthalate add the indicated number of mt of 0.2 N HCl and dilute to 200 mt Indicators thymol blue and brom phenol blue

pН	HCI	pH	ncı	pli	IICI	pH	HCI
2 2 2 4 2 6	46 ⁻⁰ 39 60 32 95		26 42 20 32	3 2 3 4	14 °0 9 90	3 6 3 8	2 63

Group 3 To ,0 ml of 0.2 M and potassum phthalate add the indicated number of ml of 0.2 \ \aOH and dilute to 200 ml Indicators brom phenol blue brom cresol green and brom cresol purple

pH	NOH	ΡΉ	`\aOH	pЦ	\aOH	pН	NaOH
4 0	0 40	4 6	12 15	5 2	29 95	5 8	43 00
4 2	3 70	4 8	17 10	5 4	35 45	6 0	45 45
4 4	7 50	5 0	23 85	5 6	39 85	6 2	47 00

Group 4 To 00 ml of 0.2 M and potassium phosphate add the indicated number of ml of 0.2 N NaOH and dulyte to 200 ml Indicators brom cresol purple brom thymol blue and phenol red

pH	NaOH	pН	MonA	рĦ	NaOH	pH	NaOH
5 8	3 72	6 4	12 60	7 0	29 63	7 6	42 80
6 0	5 70	6 6	17 80	7 2	35 00	7 8	15 20
6 2	8 60	6 8	23 65	7 4	39 50	8 0	46 80

Group 5 To 50 ml of 0.2 M born and in 0.2 M KCl add the indicated number of ml of 0.2 N NaOH and dilute to 200 ml Indicators crossol red and thy mol blue

pH	NaOH	pН	NaOH	рĦ	NnOH	pН	NaOH
7 8	2 61	8 4	8 50	9 0	21 30	9 6	36 85
8 0	3 97	8 6	12 00	9 2	26 70	9 8	40 80
8 2	5 90	8 8	16 30	9 4	32 00	10 0	43 90

Theory of Indicators, Indicators may usually be regarded as weak organic acids (or bases) whose un-ionized molecules exhibit one color whereas their amons (or cations) possess a different color. In the case of an indicator which behaves as a weak acid, for example, we have the following equihbrium reaction.

Applying the law of mass action to this ionization of the indicator acid, we get

$$\frac{[H^+][Indicator]}{[H][Indicator]} = K', \text{ or } \frac{[H^+]}{K'} = \frac{[H][Indicator]}{[Indicator]}$$

where K' is called the apparent dissociation constant of the indicator

The color imparted to a solution by the indicator depends upon the relative proportions of the two forms of the indicator present in the solution, and this, in turn, depends upon the ratio $[H^+]/K'$. Since K' is a fixed number, specific for each indicator, the color formed will depend upon the hydrogen ion concentration of the solution. In the case of brom cresol green, for example, $K' = 10^{-4}$, the undissociated molecules are yellow while the amons are blue. For this indicator, therefore, the above equation may be written

$$\frac{[H^+]}{10^{-4}} = \frac{[Yellow molecules]}{[Blue ions]}$$

If brom eresol green is added to a buffer solution whose hydrogen ion concentration is 10^{-47} (i.e., is numerically equal to the value of K' for the indicator), then according to the above equation the concentrations of yellow molecules and blue ions in the solution must be equal. In this case therefore the solution will assume the green color which results when equal numbers of these yellow and blue particles are mixed together. If, on the other hand, the indicator is added to a series of solutions whose

hydrogen ion concentrations are progressively greater than 10-47, then as the hydrogen ion concentrations increase the proportion of yellow molecules to blue ions in these mixtures will also increase Tinally at a hydrogen ion concentration of about 10 25 (pH = 38) the eye can no longer detect the presence of blue ions in the mixture. At this pH the solution exhibits the yellow color of the undissociated indicator molecules and further increases in hydrogen ion concentration will not produce any perceptible changes in color Similarly in solutions whose hydrogen ion concentrations are less than 10-47 the blue amons predominate and as the hydrogen ion concentration decreases a point is reached where it is the yellow molecules which no longer can be detected in the mixture At this pH (approximately 54) the solution exhibits the blue color of the indicator ions and further reduction in hydrogen ion concentration produces no perceptible change in the color Thus in solutions having a pH between 38 and 54 the indicator exists as a mixture of yellow molecules and blue ions and the relative amounts of each of these substances in the mixture and the colors formed vary with the hydrogen ion concentra tions of the solutions. In this range therefore brom cresol green can be used to determine the pH of an unknown solution by comparing the colo formed with the colors produced when the indicator is added to a serie of standard buffer solutions of known pH In solutions whose pH is les than 38 or more than 54 however the eye can detect the presence (only one form of the indicator and variations in pH are no longer accompanied by visible changes in color. The pH of such solutions, therefore cannot be determined by means of this particular indicator. Since eac and cator has its own value for K it follows that for each indicator the is a definite range of hydrogen ion concentration in which the indicate exists as a mixture of the ionized and un ionized forms and in which variations in hydrogen ion concentration will be accompanied by visib changes in color The problem of determining the pH of an unknow solution by means of indicators depends therefore upon the selection of an indicator whose effective range includes the pH of that particul solution

Uses of Indicators Both the concentration of free hydrogen ions if solution and its titratable acidity (or alkalinity)-i e the total amou of hydrogen (or hydroxyl) ions which the solution is capable of yields on complete dissociation-may be determined by the use of proper in cators For the latter determination the solution is titrated with stands alkalı or acıd ın the presence of an indicator which serves as an index the end point of the reaction. The indicator used for this purpose should be one which gives a sharp color change when an equivalent amount of standard acid or alkali has been added. If at this point the reaction the solution is practically neutral (pH 7) an indicator changing co at al out this reaction (litmus or rosolic acid) would be suitable. On other hand if we add 20 ml of 0 1 h AaOH to 20 ml of 0 1 h acetic stl e resulting solution will be not neutral but rather slightly alkaline due hydrolys s The sodium acetate formed will react with the water to I duce some \aOII and an equivalent amount of acctic acid Acetic : is a weak acid and will dissociate few hydrogen ions but NaOH strong base and dissociates completely, liberating OH ions which, being in excess, cause the solution to be alkaline. In titrating weak acids such as acetic and most other organic acids it is therefore necessary to use an indicator which changes color in a slightly alkaline medium. Phenolphthalein has been found to be suitable. For similar reasons the titration of weak bases such as ammonia necessitates the use of an indicator which changes color in an acid medium. Methyl red and alizarin red are often used in such cases. If a strong base such as NaOH is titrated with a strong acid such as HCl, almost any common indicator may be used, because one drop of 0.1 N solutions of these will throw the hydrogen-ion concentration so far beyond that of neutrality as to pass the turning point of any of these indicators. (For further discussion see electrometric titration method, p. 50, and titration curves, Fig. 11, p. 50.)

In making use of indicators for determination of hydrogen-ion concentration, the first problem is to choose an indicator whose effective range includes the pH of the inknown solution. That is, the indicator used must be one which, when added to the unknown solution, will exist as a mixture of both the ionized and un ionized forms, so that the color will be characteristic of the pH After selection of the proper indicator the unknown is treated with a measured volume of this indicator and the color obtained is compared with those produced when the same amount of indicator is added to a series of huffer solutions of known hydrogen-ion concentration The pH of the unknown is considered to be the same as that of the buffer solution which yields exactly the same shado of color. For use in the determination of the hydrogen-ion concentration of a solution an indicator is selected which shows a satisfactory gradation in color through the range in question, which is not readily affected by the presence of neutral salts or other substances likely to be present, and the color of which does not fade too rapidly. The ranges through which a number of indicators may be used with satisfactory results for the determination of hydrogen ion concentrations are indicated in Fig. 6. The sulforphthalem series of Clark and Lubs is especially brilliant and reliable

EXPERIMENTS ON DETERMINATION OF HYDROGEN-ION CONCENTRATION

1 Colorimetric Determination of Hydrogen-ion Concentration The first step in this procedure is to determine the approximate pH of the unknown solution so that a suitable indicator may be selected Treat a small (about a 1-ml) portion of the unknown solution with 1 drop of an indicator¹¹ solution,

If Indicators may be obtained in powder form or in prepared solutions from La Motto Chemical Freducts Co. W. A. Taylor and Co. or Hy non. Westect and Dunning all of Baltimore. Md. 0.04 per cent solutions of all of the Clark Lubs series of indicators are used. For many purposes 0.04 per cent solutions of the indicators in 0.5 per cent alcohol may be used. Clark and Lubs prefer to use aqueous solutions of the alkali saits. To prepare it exictly a stock solutions rub 0.1 g. portions of dry powder to an agate mortax with the following quantities of 0.0 a. No.01: phero if red 5.7 ml. broin phenol blue 3.2 ml. broin cresol green 2.9 ml. chlor phenol red 4.7 ml. of the capual series of the stock of the more readily obtained by using 1.1 equivalents of No.01! When solution is complete dilute to 2.5 ml. with water. Dilute portions of the except of these times to set 0.04 per cent solutions. Other indicator solutions may be made up as follows. thrains elsow 1.6 for incohence-accosalies of the contraction of the proposition of the pro

and compare the color obtained with those resulting when i drop of the indicator is added to the same volume of 0.1 N HCl (acid color) and 0.1 N NaOli (alkalne color). If the color obtained with the unknown is intermediate between the acid and alkaline colors of the indicator, the pli of the unknown lies within the effective range of this indicator and it may he used for the colorimetric determination of pli as described below. If on the other hand the unknown shows either the full acid or full alkaline color with the indicator selected, it is unsultable and another indicator must be tried in a similar manner, until an indicator has been found whose effective range includes the old of the unknown solution.

When the proper indicator has been selected, look up the effective pli range of this indicator (see chart, p. 34, or table, p. 381). Into a series of clean test tubes of clear (preferably pyrer) glass of uniform internal diameter (nbout 15 mm.), measure 10-ml. portions of standard buffer solutions covering this range, preferably in steps of 0.2 pli unit. Add 10 drops (0.5 ml.) of indicator solution to each tube and mix. Prepare a 10-ml. portion of the unknown solution in a test tube similar to that containing the standards and with the same amount of added indicator. Compare the unknown against the standards, using a comparator block (Fig. 7). The pil of the standard most closely matching the unknown is the pil of the unknown. It should be possible to interpolate between standards, thus attaining a precision of 0.1 pil unit, if the standards differ by 0.2 pil.

For approximate orientation in the preliminary trials, suitable use may be made of the ordinary laboratory indicators (litmus, phenolphthalein, methyl orange-see Fig. 6), but the final choice of indicator should be based upon trial with those indicators which have been specially selected for use in the colorimetric determination of hydrogen-ion concentration, in this connection it is well to remember that thymol blue has two effective pil ranges (1.2 to 2.8, and 8.2 to 9.8), the alkaline color for the first range as well as the acid color for the second range being obtained with some solution baving a pli between 3 and 8, such as dilute acid potassium phosphate solution. In measuring out the indicators the dropping pipet should be beld perpendicularly so that drops of constant size are obtained. In measuring out buffer solutions with a pipet the latter should not be blown out and if used for another solution must first be rinsed very thoroughly with distilled water from a flask which has not been blown through (because of the CO2 in expired air)." Color comparisons should be made in a good light using a sheet of white paper as a background.

If the stock solutions of Sørensen are used, 10-ml. portions of standard solutions may be made up from these as required. The stock solutions are measured from burets and the proportions required for a certain JH may be found in Fig. 6. Let us suppose that the unknown solution has been found to have a pil within the range of phenol red (6 to 8.2). This range is covered by the phosphate buffer mixtures. Measure out 5 ml, of the acid phosphate and 5 ml, of the akishine phosphate, mix, and add 10 drops of indicator. If the same tint is obtained as with the unknown, consult the chart. It will be found that this phosphate mixture has a pil of 6 8 and the pil of the unknown must also be 6.8 If the same tint is not obtained on the first attempt make other trials

cent aquecus solutions. Methyl red. saturated solution in 50 per cent alcohol. Neutral red. Oll per cent in 50 per cent alcohol. P-utrophend, 604 per cent in 6 per cent alcohol. Thenolphthalein 6 5 per cent alcohol. Asolution in aquecus solution. 31 For the standards it are

using proportions of 4 6, 3 7, etc. For other ranges of pll use other indicators and other buffer solutions in the same manner

If the unknown solution possesses a color of its own either of two proce dures may be resorted to The first method is to add to the standard, before any indicator is added, enough mert coloring matter to match that in the unknown and then add indicator to both 29 A better method which may also be applied to moderately turbid solutions was succested by Walpole The comparator shown in Fig. 7 is used 20 The colored fluld plus indicator is placed in a test tube in hole 3 and the standard solutions for comparison on each side in holes 1 and 5 Then opposite the standards in holes 2 and 6 are placed

tubes containing the colored unknown solution without added indicator and In hole 4 a tube containing water only Comparison is made through the apertures A. B. and C

2 Colorimetric Determination of Hydro gen-ion Concentration Without Use of Buffer Solutions at This procedure is most simply carried out using indicators changing from colorless to a single colored form such as phenolphthalein or the nitro phenols. Within the effective cance of the Indicator, the death of color varies with the pil and depends upon the amount of dis sociated indicator present The amount of completely dissociated indicator required

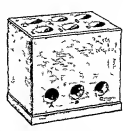


FIG 7 COMPARATOR BLOCK

to give the same tint as the unknown is a measure of the pH of the latter The standards are made up by adding measured amounts of indicator solution to portions of solution in which the indicator gives its maximum color (compiete dissociation)

To 10 ml of unknown solution in a test tube add 1 ml of indicator solution " To another tube (the standard) add 9 ml of a solution giving the

¹¹ Michaelis suggests the following indicators

Vame	Composition	pK Range 18° C of pH		Stock Solution of In licator	
8 Dinitrophenol a-Dinitropl enol by Dinitropl et ol postrop het ol m stropl enol Phenoly btbalen	1-oxy 2 G-dunitrobenzol 1-oxy 2 4-dinitrobenzol 1-oxy 2 5-dinitrobenzol	4 06	2 2-4 0 2 5-4 5 4 0 5 5 5 2 7 0 C 7 9 1 8 5-10 5	0 1 g 300 ml H ₂ 0 0 1 g 200 ml H ₂ 0 0 1 g -00 ml H ₃ 0 0 1 g 100 ml H ₃ 0 0 3 g 100 ml H ₃ 0 0 0 f g 30 ml slco- hol + 70 nl H ₃ 0	

²⁹ Solutions statable for this purpose include Bismarck brown tropacolin O and tropacolin OO in 0 02 per cent aqueous solutions cotton blue 0 01 per cent methyl violet 0 002 per cent methyl orange 0 01 per cent all in water curcumine 0 02 per cent in 60 per cent alcohol and helianthin II 0 01 per cent in 80 per cent alcohol

³⁰ A comparator may be made by boring 6 holes in purs in a block of soft wood tile paired holes being as close together as possible and just large enough to hold ordinary test tubes Smaller boles are then bored perpendicular to these Stain black with an alcol of wood stun A sheet of ground glass placed between the light source and the tubes is helpful 21 Michaelis and Gremant Brochem Z 109 165 (1920)

maximum color with the indicator (e.g., 0.02 N NaOII) and then add indicator solution diluted ten times from a microburet or Mohr pipet until a color is obtained which on diluting the standard to the same volume as the unknown (11.0 ml.) matches that of the unknown. If the first standard does not exactly match, make a second one; if more than 2 ml. of diluted indicator are needed, reneat, using undiluted indicator.

CALCULATION.

$$pH = pK' + \log \frac{c}{1 - C}$$

where pK' is a constant for the indicator (consult table) and C is the volume in mof indicator added to the standard, expressed on an undiluted basis. If 1 ml of diluted indicator were used (0.1 ml of undiluted) and the indicator is p-nitrophenol (pK' 7.18)

- 10)) = 7.18 - 0.95 = 6.23. A chart giving values of $\log \frac{C}{1-C}$ is convenient in making calculations 22

Other Colorimetric Methods for Determining Hydrogen-ion Concentration. The method without huffers may also be applied to two-color indicators (Gillespie), in this case each standard consists of two tubes set one behind the other in the Walpole comparator. One tube contains for example a known amount of phenol red in alkaline solution (red) and the other in acid solution (yellow). By varying the proportions any shade of this indicator may be obtained. For further details, see Clark.

indicator paper ("liydrion," "Accutint," etc.) may also be used for the determination of pli. Such paper is convenient and sufficiently accurate for many purposes.

3. Comparison of Hydrogen-ion Concentration and Titrotable Acidity. (A) Determine colorimetrically the hydrogen-lon concentration of a 0.01 N solution of hydrochloric acid using thymol hlue as an indicator and of a 0.01 N acetic acid solution using hrom cresol green as an indicator. Note the great difference between the true scidities of the two solutions,

Titrate 10-ml. portions of 0.01 N hydrochloric acid and of 0.01 N acetic acid with 0.01 N NaOli, using phenolphthalein as an indicator. Note that identical results are obtained for the titratable acidities of the two.

(B) Mix equal portions of M/15 potassium dlhydrogen phosphate and M/15 disodium phosphate (see chart). Note that the mixture is practically neutral to iltmus. Titrate one 10-ml, portion of this mixture with 0.1 N NaOil, using phenoiphthalein as an indicator. Titrate another portion with 0.1 N HCi solution, using methy i orange. Explain results.

ELECTROMETRIC DETERMINATION OF HYDROGEN-ION CONCENTRATION

Electrode Potentials. If an excess of solid glucose is placed in water, some of the molecules of the solid, being in a continual state of vibration, escape into the liquid. As the molecules of glucose in solution accumulate, some of them collide with the particles of solid and are retained. An equilibrium is finally established in which there is a balanced exchange of molecules between the solid and the solution. At this stage the solution is saturated with glucose and the solution pressure of the solid is said to he equal to the osmotic pressure of the dissolved substance.

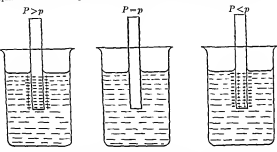


Fig. 8. Deagram of Relations at Suppace of a Metal Dipping into Solutions Containing Its lons in Different Concentrations (P = solution pressure of the media. p = sometic pressure of metallic ions.)

If a strip of metal (electrode) is dipped into water, it also tends to dissolve owing to its solution pressure, P. The metal goes into solution as positive ions leaving its electrons (presumably less firmly bound valence electrons) behind as a negative charge on the electrode. This charge, by attracting the positive ions of metal already in solution, builds up an electrical double layer at the interface between electrode and solution which opposes the entrance of any more positive ions into the solution, and the process of solution ceases therefore when this charge becomes large enough to prevent the further separation of positive ions from the metal. Although equilibrium is established before a measurable quantity of the metal dissolves, since the charge on each ion is comparatively large, a measurable potential difference is developed between the electrode and the solution.

When a metal is dipped into a solution of one of its salts the conditions are somewhat altered because the positive ions of the metal already in solution oppose the separation of more ions from the metal. The point of equilibrium therefore will depend upon the relative values of the two opposing forces; the solution pressure (P) of the metal and the osmotic pressure (p) of its ions in solution. We may therefore distinguish three possibilities (see Fig. 8);

1 P > p In this case the metal continues to send positive ions into the solution until the accumulated charges oppose further action The metal thus acquires a negative charge relative to the solution

2 P < p In this case the positive ions in solution deposit on the metal yielding to it their positive charges, until the accumulated charges oppose further deposition. The metal thus acquires a positive charge relative to the solution

3 P = p In this case there is neither solution nor deposition and no

potential difference develops between the metal and the solution

Vernst has shown by thermodynamic reasoning that the potential difference between a metal and a solution of one of its salts is given by

$$E = \frac{RT}{nF} \ln \frac{P}{p}$$

where E = electrode potential R = gas constant = 8316 joules per degree, T = absolute temperature, n = valency of metal ion F = farada) = 90 500 coulombs P = solution pressure, and p = osmotic pressure of metal ions in solution (The symbol "In" stands for the natural logarithm)

I rom this equation it becomes evident that, since RT/F is a constant for any given temperature the magnitude and sign of the electrode poten tial F, is determined by n and P, which depend solely upon the nature of the metal used, and p which is a function of the concentration of metal ions in solution. If therefore we have two electrodes of the same metal dipping into solutions of different concentrations the two electrode poten tials will be different. And if we make suitable connections24 between the two electrodes and the two solutions an electrical current will pass from ons electrode to the other, the electromotive force of which will be equal to the difference between the two electrode potentials-1 e

$$\begin{split} F &= E_1 - E_2 \\ &= \frac{RT}{n\bar{F}} \ln \frac{P_1}{p_1} - \frac{RT}{n\bar{F}} \ln \frac{P_2}{p_2} \\ &= \frac{RT}{n\bar{I}} \left(\ln P_1 - \ln p_1 - \ln P_2 + \ln p_2 \right) \\ &= \frac{RT}{n\bar{I}} \ln \frac{p_2}{p_1} \left(\text{since } P_1 = P_2 \text{ for the same metal} \right) \end{split}$$

If esmotic pressure is regarded as proportional to concentration, we may replace pr and pr by cr and cr and get

$$L = \frac{RT}{nF} \ln \frac{c_2}{c_1}$$

An arrangement such as here described is known as a concentration cell, and the above equation permits us to calculate the concentration of metal ions in an unknown solution from the electromotive force developed when that solution is combined in a cell with a similar solution of known ionic concentration. It should be noted that the concentration calculated by the use of the above equation will be that of electromotively active ions, which is not necessarily the same as that obtained by ion conductivity or other means. For a further discussion of this point, the reader is referred to Clark.

to Clark The Hydrogen Electrode. A strip of platinum coated with platinum black and saturated with hydrogen gas, acts exactly as the metal electrodes described above when dipped into a solution containing hydrogen ions. We can therefore construct a concentration cell by dipping two such cleetrodes into two solutions of different hydrogen ion concentrations. and, if the concentration of hydrogen ions in one solution is kept at a known constant value we can use the electromotive force developed by such a cell to calculate the hydrogen-ion concentration in the second unknown solution. If the concentration of hydrogen ions in the known solution is normal the equation for the electromotive force becomes $E = \frac{RT}{nF} \ln \frac{1}{(H^+)}$ By substituting the values for R, T n, and F at 25° C and multiplying by 2 303 to change from the natural to the common system of logarithms, we get $E = 0.059 \log \frac{1}{(H+1)}$ or E = 0.059 (- log [H+]) Hydrogen ion concentrations following the convention intro duced by Sørensen, are usually expressed in terms of pH by definition, $pH = -\log [H^+]$ Therefore, E = 0.059 pH or $pH = \frac{E}{0.059}$

In practice it is convenient to use, instead of a hydrogen electrode in a normal solution of hydrogen ions, a calomel electrode which has been standardized against such a solution. A calomel electrode is made up of mercury in contact with calomel in a solution of potassium chloride, the latter being either normal, tenth normal, or, more often, saturated. The normal hydrogen electrode is approximately 0.25 volt more positive than the saturated calomel electrode is that the difference of potential between the calomel electrode and the unknown will be 0.25 volt greater than that between the latter and the normal hydrogen electrode. Therefore, using the saturated calomel electrode at a temperature of 25° C, pH = $\frac{E-0.246}{0.059}$ We may read E directly in volts on a voltmeter, or by

pH = $\frac{1}{0.059}$ We may read E directly in volts on a voltmeter, or by me ins of a potentiometer, so that the calculation becomes very simple Variations of pH with E may be plotted in the form of a curve. In some

instruments the voltmeter is graduated to read pH directly

Regardless of the types of electrodes used in the chain, the measurement of E must be conducted by means which do not entail an appreciable flow of current from the system, because such a flow introduces polyment of current from the system.

¹³ This figure varies with the temperature and with the concentration of hCl solution in the calomel electrode (see values for Le in the table on p. 45)

effects at the electrodes. The potentiometric method is ideally adapted for this purpose. The principles involved in this method are described in detail in Clark's book, and also in the descriptive literature which accompanies the special apparatus required. 25

Hildchrand's method described below, requires only such electric apparatus as is usually found in any laboratory, and serves to illustrate

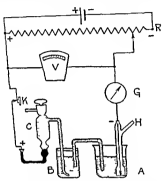


FIG 9 ABRANCIMINT OF APPARATUS FOR I LECTRO-METRIC DETERMINATION OF HYDROGEN ION CONCEN-TRATION (METHOD OF HILDRIDGEN))

H is the hydrogen electrode C the caloniel electrode S a solimeter G a galvanometer R a rheostat A the beaker containing the unknown solution and B are innerting vessel

the principles of the electrometric method. The use of a potentiom instead of the coltmeter indicated increases the accuracy of the deternation and is to be preferred.

Determination of Hydrogen ion Concentration by Hildebrand's Meth Set up the apparatus as indicated in Fig. 9. The battery may be an ord dry cell. The voltmeter 3 should have a range of 12 volts and scale division 800 volt. The portable galvanormeter G should have a sensitivity of the properties of t

the calomel electrode ²⁸ H is a hydrogen electrode of the Hildebrand type especiolly suitable for solutions not containing protein The Clark electrode is a type frequently employed in biochemical work and may be used with a shaking device to ossist in obtaining equilibrium rapidly.

The essential part of the hydrogen electrode is the piece of plotinum foil which is to dip into the unknown solution. This must be given a spongy coot of plotinum black so that it will retain as much hydrogen as possible and

must be saturated with hydrogen just before use 29

The unknown solution is placed in beaker A. The calomel electrode is clamped in place, os olso is the prepared hydrogen electrode A stream of hydrogen bear solutions as side arm of the hydrogen electrode so that it bubbles up through the solution of the rate of about two bubbles per second Beaker B contains saturated KCl solution. The connecting tube contains saturated KCl made up in warm 3 per cent agar and ollowed to solidify in the tube. This prevents syphoning between B and A.

Stir the solution well on automatic stirring device is best for this purpose Move the silding contact on the rheostat until the voltage drawn from the dry cell is equal to thot produced in unknown solution. This is indicated by the fact that no current passes through the galvanometer when the contact key K is pressed down, and there is therefore no deflection of the needle. Until a balance is reached the spring contact key should be closed only momentarily by a slight tap. At the balance point read the voltmeter.

CALCULATION OF pH See the discussion preceding Determination At 25° using

CALCULATION OF pH. See the discussion preceding Determination At 25° using saturated calomel electrode pH = (E - 0.246)/0.059 If for example the voltmeter reads 0.652 then pH = (0.652 - 0.246)/0.059 = 7.0 and the solution possesses a neutral reaction Corrections for different electrodes and temperatures are gn eo in the following table. This apparatus should have an accuracy of about 0.1 pH. Test the apparatus first, using buffer solutions of known pH. 4°

"Hisdogen may be purchased in cylinders or may be prepared by electrolysis of an AaOH solution or from pure zinc. To purify pass through a wash bottle containing alkaline lyrogallol solution and two britles containing water. Connect one of these bottles with a third tube dip ping below the surface of innerury in another vessel. This acts as a safety

With gas tanks a gas regulator for adjusting flow is desirable

[&]quot;Calomel Electrode Place about 3 mi of carefully purified mercury in the bottom of the clectrode vessel Rub together mercury and mercurous chloride (calomel) with a little sat urated KCI solution to form a paste Place a layer of this paste about a half into thick over the mercury and fill the vessel to the end of the side tube with anturated kCI solution II it is desired to make a normal or tenth normal calomel electrode corresponding solutions of kCI should be used instead of the saturated solution II is convenient to make a permanent connection between the upper side arm and a reservoir bottle containing KCI solution B; running a little of the solution through the calomel cell before each series of determinations the side arm may be washed free from maternals that may have diffused up into: The storp-cock of this electrode should be tight but should not be greased. A typical calomel electrode is shown in C Fig. 9.

[&]quot;Clean the platnum electrode thoroughly with chronic and Connect with the negative pole of a dry cell and connect another strip of platnum with the positive pole. Dip both in a dilute (2 per cent) solution of platnuc chloride Electrolyse for 10 to 15 minutes free the current Dip into 10 per cent suffure veid with the electrode as the cathode and let the current run for a few minutes. The platnum is saturated with hy drogen Wash with distilled water. Keep under distilled water when not in use. It is platnum coat should last for several weeks but must be saturated with hy drogen just before use. Instead of a strip of platnum a wire alout 1 mm in diameter slightly flattened at the end may be used. Fquilibrium with such an electrode may be brought about more rapidly. Shorter periods of electrodeposition produce electrodes which come to equilibrium more rapidly but which are not so permanent.

⁴ Suntable solution tend! normal with respect to both acetic acid and sodium acetate may be use 1 4.25% with the estimated cal smel electroded 11s gives 6.51% volt and with the 0.1 N electrode 0.61% volt and 100 ml. N acetic acid with water to make 500 ml.

STANDARD VALLES FOR CALONEL I LECTRODES (Referre I to the Normal Hydrogen Electrode)

	F . for Deff			
Ter perat re	01 Y	10 M	Saturated (Approximate Potential)	2 303 RT nF
18	0 3380	0 2864	0 2.06	0 0577
20	0 3379	0 2860	0 2492	0 0581
25	0.33-6	0 2848	0 2464	0 0591
30	0 3372	0 2836	0 2437	0 0601
40	0 3360	ì	1	ì

do this is the most frequent eause of error with a pH meter, since the calibration may change significantly from time to time, particularly if the glass electrode is new or has been allowed to become dry. It is best to keep both glass and calomel electrodes immersed in water when not in use. The temperature of the solution will also influence the determination of pH. In the Beckman pH meter illustrated (Fig. 10), for example, the temperature control must be set at the temperature of the unknown solution before the reading is made

The glass electrode is used for determinations of pH in all types of biological systems. Since (unlike the hydrogen electrode) it does not cause

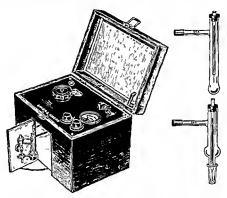


Fig. 10 Glass-Electrope Apparatus (Beckman). Glass electrode shown at right (above) and calomel electrode (below).

is to bubble through the solution, it may be used on systems like blood hich contain dissolved CO₂. The sample is not contaminated, so that ic method is especially applicable where the amount of material is mited. Electrodes are available which require only a few tenths, or even undredths, of a milliheter of solution for each determination. Determinations with the glass electrode are rapid, and with most instruments the ceuracy is within 0.05 pH. It may be used on unbuffered, colored, or arbid solutions and solutions containing sediments. With the ordinary lass electrode, determinations are accurate up to a pH of 10 or even igher, provided the solutions do not contain appreciable quantities of edium, lithium, or potassium ions. Sodium ions, especially, evert a coniderable effect on determinations in solutions with a pH of 9 or above, hen they are present in a concentration of normal or greater. Corrections may be applied in such cases (Dole). Special electrodes have been

developed for use in strongly alkaline solutions, for solids such as cheese and for many other purposes

Electrometric Titration of Acids and Bases. In many cases the use of indicators in titration is unsatisfactory because of color or turbidity of solutions to be examined or the presence of buffer substances which make the end point uncertain. The hydrogen or class electrodes may be used in such cases in titrating solutions to a definite acidity or alkalimity.

This method also makes it possible to follow the changes in acidity during the course of a titration and gives information as to the character of the

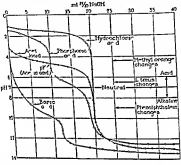


FIG. 11 TITRATION CURVES OF VARIOUS ACIDS OBTAINED BY I LYCTRO-MITTER MYSBER REMENT

Twenty ml of 0.1 \ge 1 y free lone an 1 acetic ac 1s used. I hosphoric acid about z_3 molar Horic acid saturated solution

acids or bases in solution from applications of this method are illustrated in Fig. 11 $\,$

Procedure The apparatus already described and illustrated in Fig. 9 may be used The unknown solution is placed in heaker A, into which are introduced both the hydrogen and calomel electrodes. Beaker is and the connecting tube are omitted. Hydrogen is hubbled through the solution and acid or alkall added from a hurer with stirring. At intervals read the volumeter Flor a curve of voltage or pil against milliliters of acid or alkall added. Neutralization is in licated by a rapid change of voltage or pil I Di or tribasic acids or bases may show two or more such changes. See the curve for titration of phosphoric acid. The

The glass electrode pil meter is particularly suitable for this purpose Special electrodes with long leads are available for direct titration in all ordinary beaker

in [H+] at the end point is very sharp because, as long as even 0.1 ml. of the acid remains unneutralized, the pH is still quite low and 0.1 ml. of 0.1 N NaOH in excess gives a very large increase of pH. Evidently either methyl orange, litmus, or phenolphthalein may be used in the titration. With acetic acid however the beginning [H+] is lower because this acid is weakly dissociated. The change in pH is at first fairly rapid because the sodium acetate formed yields acetate ions which repress the ionization of the acid. As the titration is continued the ionization of the acid is almost completely repressed by the larger amounts of acetate formed. When an equivalent amount of alkali has been added, the reaction is alkaline owing to slight hydrolysis of the sodium acetate to form a cetic acid (nearly all undissociated) and highly dissociated NaOH. The actual end point of the titration is therefore on the alkaline side and phenolphthalein is the proper indicator.

Fig. 11 shows also how it is possible to titrate the first hydrogen of phosphoric acid using methyl orange as an indicator and the second using phenolphthalein; the third cannot be titrated hecause the change is too gradual. Note also that boric acid cannot be titrated with phenolphthalein as an indicator.

Electrometric Oxidation and Reduction Titrations. When an ion is oxidized it loses electrons or negative charges; when it is reduced it gams electrons. Thus when Fe⁺⁺ is oxidized to Fe⁺⁺⁺ there is a loss of one electron. The process is therefore similar to that resulting in the loss of an electron when hydrogen passes into solution as H⁺. If a platinum wire is placed in a solution containing ferrous and ferric ions, the ferrous ions tend to give up electrons to the platinum and the ferric ions tend to take them away. A difference of potential will arise depending upon the concentrations of the two ions. If an oxidizing agent is now added there will he a change of potential, slow at first and then very rapid, as the concentration of ferrous ions becomes very small compared to that of the ferric ions. This rapid change of potential marks the end point of the titration.

Procedure. The apparatus used is the same as for the electrometric thration of acids and bases. The platinum electrode should however be small and should not be platinized but be bright, and kept in 1:1 HGl when not in use. The calomel electrode should be connected to the negative side of the main circuit inatead of the positive. Instead of a beaker a flask may be used and a current of CO₂ passed over the surface of the solution (not through it) to prevent oxidation by the air. "Run in an oxidizing or reducing agent from a buret. Readings of the voltimeter are taken at intervals and at the end point a drop or two of solution produces a marked change of potential.

An electronic pli meter with the glass electrode replaced by a platinum electrode may also be used. See the manufacturer's directions for the proper connection and use.

The Polarograph. Instead of adding a reactant to the solution and using the change in potential at an electrode for quantitative purposes as

⁴³ If desired the voltmeter may be eliminated. In this case adjust the rheostat so that there is no galvamoneter deflection at the beginning of the tilration. The end point is indicated by the galvamoneter needle being thrown off the scale.

described above it is possible to apply an external voltage across the electrodes and electroly ze the ions by foreing them to react at the electrodes At the cathode for example the univalent metallic ion M⁺ will accept a electron ϵ and be reduced to the metallic state

$$M^+ + \epsilon = M$$

Under the proper conditions the voltage required to bring about such a raction is characteristic of the chemical nature of the ion and the curre resulting from electron transfer may be used as a measure of the concetration of the ion

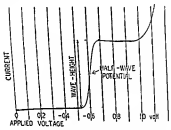


Fig. 12 Typical Polarogram

Air free solution of 0.001 M CdSO₄ in 0.1 M CoCl₁
buffered at pH 5

From Müller J Chem Education 18 71 (1941)

The instrument employed to apply a known and variable voltage across the electrodes and to measure the resulting current is called the polarograph (Heyrovsky) A simple form of polarograph which can be assembled from equipment available in most laboratories has been described by Müllert 'I he graphical recording of the relation between applied voltage and current is called a polarogram (Fig. 12). Most commercial polarographs 'I provide for the continuous and automatic recording of the polarogram using automatic recorders or photography.

In its simplest form the current-voltage curve for a particular ion is approximately symmetrical around the half wave potential which is determined by the kind of ion undergoing reduction. Under the proper conditions the wave height becomes a measure of the concentration of the ion. The concentration of material required for polarographic analysis is quite low ranging from 10. M to 10. M under ordinary circumstances in a volume of 1 ml or more. It is worthy of note that solutions may be re-

Carbohydrates

Definition. The name carbohydrates is given to a class of substances which are especially prominent constituents of plants and are found also in the animal body, either free or as components of certain proteins, lipides, and other compounds. They are called carboliydrates because they contain the elements C, H, and O, the H and O heiug present in the proportion to form water. The term is not strictly appropriate inasmuch as there are compounds (such as acetic acid, lactic acid, and inositol) which have H and O present in the proportion to form water, but which are not carbohydrates, and there are also true carbohydrates which do not have H and O present in this proportion—e.g., rhamnose, C6H12O5, Chemically considered the carhohydrates are aldebyde or ketone derivatives of polyhydric alcohols, or condensation products of such substances. The aldehyde derivatives are spoken of as aldoses, and the ketone derivatives are spoken of as ketoscs. It is worthy of note that among pure organic compounds, the one prepared in the largest quantity in the United States is sucrose, a carhohydrate.

Classification. The carhohydrates are usually classified, according to the number of simple carbohydrate groups which they contain, as mono, di-, tri-, and polysaccharides. The mouosaccharides, which contain only a single such group and cannot therefore he hydrolyzed into simpler substances, are further characterized according to the length of the carbon chain as trioses (CaHoO), teroses (CaHoO), pentoses (CaHoO), twooses (CaHoO), etc. The disaccharides give two molecules of simple sugars on hydrolysis and the more common ones have the general formula CaHoO). The polysaccharides give many molecules of simple sugars on lydrolysis and the formula for the more important members of this group is (CaHoO), where x represents the number of simple sugar groups present. In a general way the soluhility of the carbohydrates varies with the complexity, the more complex being the less soluble. This means simply that, as a class, the monosaccharides (heaoses) are the most soluhle and the polysaccharides (starehes and cellulose) are the least soluhle.

The more common carbohydrates may be classified as follows:

- Monosaccharides.
 - Hexoses (C₅H₁₂O₅). Glucose, fructose, galactose, mannose.
 Pentoses (C₅H₁₀O₅). Arabinose, xylose, ribose, rhamnose
 - (methyl-pentose) C₆H₁₂O₆, deoxyrihose (C₅H₁₀O₆).

II. Disaccharides (C1-H2:O11).

Maltose, lactose, sucrose, gentiobiose, isomaltose, cellohiosc.

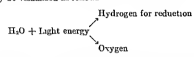
- III Trisaccharides (C18H22O16)
 - Raffinose
 - IV Polysaccharides (Cell10Os)z
 - 1 Starch Group Starch dextrin, glycogen, mulin
 - 2 Cellulose Group
 - (a) Cellulose
 - (b) Hemicelluloses
 - (1) Pentosans Gum arabic
 - (2) Hexosans Galactans, agar-agar, dextrans, levans
 - (3) Hexo-pentosans Pectin

Photosynthesis. It is interesting to note that the bulk of these highly complex carbohydrates found in nature, whose structures in many in stances are as yet unknown, are synthesized by plants from such simple precursors as water and carbon dioxide. Although the mechanism of such synthesis is still obscure, it has long been recognized that the synthetic process is related to the presence of sunlight and chlorophyll or other such coloring matter in the plant.

It is generally agreed that photosynthesis occurs in two distinct phases, one involving reactions which proceed in the absence of light (dark reaction or Blaci man reaction), and tho other involving the presence of light and a photosensitive catalyst such as chlorophyll or other pigment Opinions vary as to the relative contributions of these two phases to the total process, earlier views postulated that carbon dioxide utilization took place in the dark, but oxygen evolution was associated with the presence of light and chlorophyll. More recent views incline to the belief that hoth carbon dioxide assimilation and oxygen production are dark reactions in the sense that they do not fundamentally require light, the function of the light being to initiate and maintain the sequence of dark reactions by the photochemical decomposition of a suitable substance, possibly water itself (see below), in the presence of the photocatalyst chlorophyll or similar pigment.

The simplest possible formulation of the over all reaction for the production of a carbohydrate such as glucose (C₄H₁₂O₄) from carbon dioxide and water by photosynthesis is the following

by the use of the oxygen isotope O¹⁸ (heavy oxygen, see p. 983) as a tracer indicates that the oxygen evolved during photosynthesis is derived from the water entering the reaction and not from the earbon dioxide utilized Furthermore, the oxygen of the water that is formed is derived from the earbon dioxide. Hence the reaction is essentially a reduction, water supplying both the hydrogen required for the reduction and the oxygen which is evolved. According to this view, the role of water in the photochemical process may be visualized as follows.



The availability of earbon dioxide labeled with radioactive earbon has stimulated research into the mechanisms of earbon dioxide assimilation during photosynthesis. The results of Calvin and his associates are of particular interest in this councetion. Using various green plants (grains, algae), these investigators were able to show that, following a brief period of photosynthesis in the presence of radioactive earbon dioxide, a number of compounds containing radioactive earbon dioxide, a number of compounds containing radioactive earbon dioxide from the plant extracts and identified Paper chromatography (see p. 16) proved to be of particular value in these experiments. As the period of photosynthesis was made progressively shorter until it was reduced to a matter of only a few seconds' exposure to light, the number of different labeled compounds isolated was reduced until only one substance stood out as a major bearer of the assimilated radiocarbon—this substance is phosphogly cero-acid.

CH₂OPO₃H₂
CHOH
COOH
Phosphoglyceric acid

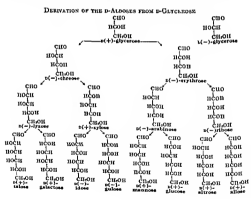
The radiocarbon is found more abundantly in the carboxyl group of the phosphoglycene and than in the other two carbon atoms, this leads to the belief that the first step in CO-inptake during photosynthesis is the condens ition of CO₂ with some 2 carbon precursor to form phosphoglycene and The nature of this precursor is not known, the finding of the isotope label in the other two carbon atoms of phosphoglycene and, but to a lesser extent, indicates that the precursor is produced by some eyelic process but to be worked out in which phosphoglycene and itself plays a pitt i tentitive cycle involving 5 carbon and 7 carbon sugars as intermediates his been proposed, but the details remain obscure.

Phosphogly cene and is a well recognized intermediate in the break-down of carbohy drute by animal tissues (see p. 275). Indeed, the student of numal brochemistry will note a striking resemblance between the reactions of plant photosynthesis thus far described and those already known for carbohy drate metabolism in animal tissues. Furthermore, CO₂-

assimilation also occurs in animal tissues to a limited but definite extent, as discussed in Chapter 33, and there appears to he much in common between the processes as they occur in plants and animals. The action of light in photosynthesis remains obscure; according to one view, the light energy acts by opening the S—S bond in a compound such as 6,8-thioetic acid (a-lippic acid, see Chapter 35). This would then he the reduction reaction mentioned above; how this fits in with the water reaction remains to be established.

It should not be inferred that photosynthesis is of importance solely in connection with cathohydrates. Plants also produce nitrogenous and non-nitrogenous substances such as ammo acids, proteins, and fats, which are of fundamental significance to animal nutrition, and there is no reason to believe that the energy of photosynthesis is not used directly or indirectly for the formation of these compounds also.

Spatial Configuration of the Sugars. The triose glycerose (also called glyceraldehyde) has one asymmetric carbon atom and therefore



light by the compound. If it is desired to indicate the direction of rotation for the compound, the symbols (+) meaning dextrorotatory and (-) meaning levorotatory, may be used. Thus a dextrorotatory compound with the p configuration may be indicated as p(+), while a levorotatory compound with the same configuration has the symbol p(-).

As the number of asymmetric carbon atoms in the sugar molecule increases on going from the trioses to the higher monosaccharides, the number of stereoisomers increases in accordance with the van't Hoff formula 2a, where n represents the number of asymmetric carbon atoms. All of these stereoisomers which have a configuration identical with that of p-glycerosc around a suitably selected reference asymmetric carbon atom are called D sugars, regardless of their direction of rotation. The reference carbon atom is, for sugars containing more than one asymmetric carbon. the asymmetric carbon atom farthest removed from the active (i.e.; aldehyde or ketone) end of the molecule. These relationships for the p aldoses are illustrated by the preceding chart, in which the reference asymmetric carbon atom is indicated by heavy black type. In this connection note that if the structure is written with the active group at the top it is conventional to write the OH group on the right-hand side of the reference asymmetric carbon atom for the p configuration and on the left-hand side for the L configuration. Naturally occurring sugars are mainly of the p configuration; L sugars have, however, been isolated from certain plant and animal sources

MONOSACCHARIDES

HEXOSES, C₆H₁₂O₆

The hexoses are monosaccharides containing a chain of six carhon atoms in the molecule. They may be either aldoses or ketoses. They are among the most important of the simple sugars and occur widely distributed in nature, either in the free state or in combination with other molecules, from which they may usually he separated by hydrolysis. The most important hexoses biologically are glucose, fructose, galactose, and mannose. Of these, fructose is a ketohexose; the others are aldohexoses The varnous aldohexoses differ structurally from one another solely in the spatial arrangement of the H and OH groups around certain of the carbon atoms in the molecule (see p. 58). This difference may result in markedly different physiological properties. As a class the hexoses are extremely soluble, are optically active, and possess certain characteristic general and specific chemical properties which are used in their identification and determination.

CHOH).

Glucose, also called dextrose or grape sugar, is found widely distributed in nature either in the free state or combined with other compounds, To-

gether with fructose, it occurs in the juice of many fruits, and is obtained commercially by the hydroly is of starch. It is the sugar found in the blood, where its concentration normally is about 0 I per cent. Ordinarily it is not present in the urine except possibly in traces, but appreciable amounts are found under certain conditions, such as in dishetes melliting.

It is dextrorotatory in solution, having a specific rotation of $+52.5^{\circ}$ As an aldohevose it may be exidized to the corresponding sugar acids or reduced to an alcohol

$$\begin{array}{c|cccc} CII_2OII & CHO & COOH & COOH \\ \downarrow & (red) & \downarrow & (ox.) & \downarrow & (ox.) & \downarrow & (ox.) \\ (CIIOII)_4 & & (CHOII)_4 & & (CIIOH)_4 & & (CHOH)_4 \\ \hline CH_2OII & CH_2OH & CH_2OH & COOH \\ Sorbitol & Glucose & Gluconic \\ & & acid & acid & acid \\ \end{array}$$

Another oxidation product, glucurome acid, CHO (CHOH), COOH is of considerable physiological importance, hung found in the urine in combination with certain exerctory products (see Chapters 20 and 29)

In freshly prepared solutions of the usual crystalline glucose the α form exists alone, the solution having a specific rotation of $+113.4^{\circ}$. On standing, however, some of the molecules are slowly transformed into the β form and an equilibrium is finally established between the two forms. A trace of alkali hastens the formation of this equilibrium. The specific rotation of the solution falls as the amount of the β form increases until equilibrium, with its characteristic specific rotation of $+52.5^{\circ}$, is reached. This phenomenon is known as mutarotation. Mutarotation is observed not only with glucose but with all sugars which have a free or potentially free aldehyde or ketone group in the molecule.

Iu addition to the amylene oxide configurations, glucose may exist in forms where the terminal carbon atom is linked through an oxygen atom to other carbon atoms than the fifth, thus forming ethylene, propylene, and hutylene oxide rings each of which may exist in the α and β forms. Of these various forms the butylene oxide form predominates, but there is evidence that all of the forms mentioned may exist in glucose solutions. These modifications are more reactive than the ordinary amylene oxide form and are presumably responsible for the concept of an "active" form of the glucose molecule, the so-called γ -glucose.

 α and β forms, respectively, of butylene oxide structure of D-glucose

Haworth has pointed out that the amylene oxide and butylene oxide rings are structurally related to pyran and furan, respectively. Normal

glucose, using the nomenclature proposed by Haworth and now widely used, thus becomes glucopyranose while the butylene oxide form is called splucofuranose.

A ray examinations indicate that the five carhon atoms of the pyranostring he in the same plane while the oxygen atom lies in a different plane. The H and OH groups are placed above and below this plane of carboratoms in accordance with the chemical evidence concerning the positions of these groups in the sugar molecule.

In common with other reducing sugars, glucose is quite labile to the action of alkali λ trace of alkali accelerates the attainment of equilibrium between the α and β forms Further contact with alkali results in the appearance of fructose and mannose in the solution in equilibrium with glucose, probably through the intermediate formation of a common conditution rinvolving the first two carbon atoms Strong alkali (with heating) decomposes the molecule completely producing a number of smaller fra₂ments and condensation products which usually give a hrown color to the solution (Moores test, now obsolete)

Glucose is one of the sweetest of the common sugars being excelled only by fructose and sucrose. The following table gives the relative sweetness of some of the sugars considering sucrose as 100 low) without the preliminary formation of an insoluble hydrazone (mannose, p. 75) and in the absence of a positive Resorcinol-Hydrochloric Acid Reaction (fructose, p. 73)

- 1. Solubility Test the solubility of glucose in water and in alcohol If in doubt about the solubility of a compound, filter from excess solid and test the filtrate for the substance in question, or if the solvent is nonaqueous, allow it to evaporate and examine it for a residue
- 2 a-Naphthol Reaction (Molsch) To 5 ml of sugar solution in a test tube, add 2 drops of Molisch's reagent (a 5 per cent solution of α -naphthol in alcobol) Mix thoroughly Incline the tube and allow about 3 ml of concentrated sulfure acid to flow down the side of the tube, thus forming a layer of acid beneath the sugar A reddish-violet zone appears at the junction between the two liquids Repeat the test, using 5 ml of 0 1 per cent furfural solution in stead of the sugar Instead of α naphthol, 3 to 4 drops of a 5 per cent alcoholic solution of thymoil may be used?

The reaction is due to the formation of furfural and furfural derivatives, such as hydrolymethylfurfural, by the acid acting on the sugar

The test is thus given by furfural and all furfural-yielding substances and is not a specific test for carbohydrates. Concentrated solutions of organic compounds may give a red instead of a violet color due to the charming action of the sulfuric acid. In case of doubt the reaction should he repeated on a more dilute solution of the material to be tested.

3 Phenylhydrazine Reaction To a small amount of phenylhydrazine mixture (about 0.5 inch in a small test tube), add 5 ml of the sugar solution, shake well, and heat on a boiling water bath for one-half to three quarters of an hour Allow the tube to cool slowly (not under the tap) and examine the crystals microscopically (see Plate II) Better crystals may be obtained if the tubes are allowed to cool in the water bath

If the solution has become too concentrated in the boiling process it will be light red in color and no crystals will separate until it is diluted with water

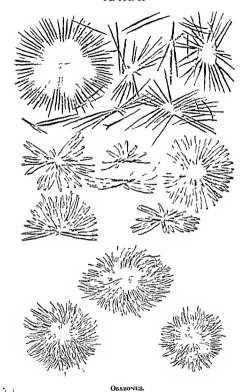
Yellow crystalline compounds called osazones are formed from certain sugars under these conditions, each individual sugar in general giving risk to an osazone of a definite crystallino form which is typical for that sugar. It is important to remember in this connection that, of the simple sugars of

¹ Thymol has the advantage that its solutions do not deteriorate Like it e a naphthol tat it is also pare by alleddydes and by acids such as forme lactic oxalic citric etc and by actione. The tests are very delicate being given by solutions of 0 001 per cent glucose and 0 0001 per cent sucross.

See Appendix.

interest in physiological chemistry, glucose, fructose, and mannose yield the same osazone because of similarities in their molecular structures of the various osazones, it is generally possible to recognize glucosazone by its crystalline form Malto-azone may also be recognized if it happens to crystallize in its most characteristic form. Otherwise it may not be possible to distinguish it from lactosazone. The melting points of the recrystallized osazones may be used as further means of identification, but since they extend over a range of several degrees and are not far apart for the different sugars, the determination of melting points is of doubtful value ⁴ The reaction with glucose is sufficiently delicate to produce under favorable conditions a visible precipitate with solutions containing as little as 0.01 per cent glucose

The reaction leading to the formation of gluco azone is indicated by the following equations: The intermediate steps between hydrazone and osazone are not known with certainty, despite earlier views to the contrary



Upper form, dextrosazone (glucosazone), central form, maltosazone, louer form, lactosazone

In testing a solution preserved by chloroform a positive reaction may be obtained in the absence of sugar. This is due to the fact that the hot alkali produces reducing substances from the chloroform.

Aminonium salts also interfere with Lehling's test. If present in excess the solution (e.g. urine) she tild be made alkaline with \a_2CO_1 and boiled in order to decompose the ammonium salts. Prolonged contact with hot strong alkali may lead to destruction of the sugar present.

If the solution under examination by I chling s test is acid in reaction it must be neutralized or made alkaline before applying the test

5 Benedict s Test Benedict modified the Felling solution to produce an Improved reagent which has largely displaced the latter in routine laboratory practice. The following is the procedure for the detection of glucose in solution To 5 ml of the reagent in a test tube add exactly 8 drops of the solution under examination Mitwell Boil the mixture vigorously for two minutes or place in boiling water for three minutes) and then allow the fluid to coll spontaneously (do not hasten cooling by immersion in cold water) in the presence of dextrose the entire body of the solution will be filled with a precipitate which may be red yellow or green in color depending upon the amount of sugar present in the presence of over 0.2 to 0.3 per cent of glucose the precipitate will form quickly if no glucose is present, the solution will remain perfectly clear

Even very small quantities of glueose (01 per cent) yield precipitates of surprising bulk with this reagent, and the positive reaction for glueose is the filling of the entire body of the solution with a precipitate, so that the solution becomes opaque Since amount rather than color of the precipitate is made the basis of this test it may be applied even for the detection of small quantities of glueose, as readily in artificial light as in daylight Chloroform does not interfere with this test nor do urre eard or creatinine interfere to such an extent as in the case of Felling 8 test.

c Binnuth Reduction Test (Nylonder) To 5 ml of sugar solution in a test tube add one tenth its volume of Nylander a reagent and heat for five min utes in a bolling water bath The solution will darken if reducing sugar is present and upon standing for a few moments of believe the superior of the solution will conserve the superior of the superior of

- (a) $B_1(OH)_2NO_3 + KOH \rightarrow B_1(OH)_3 + KNO_3$ (b) $2B_1(OH)_3 - 3O \rightarrow 2B_1 + 3H_2O$
- d. Barfoed's Test. To 5 mi of Barfoed's solution in a test tube, add 0.5 ml. of giucose solution and heat to boiling Reduction is indicated by the formation of a red precipitate of cuprous oxide. If the precipitate does not appear after boiling for 30 seconds, allow the tube to stand for about 15 minutes and example.

To compare reactions of mono- and disaccharides, piace 0.5-mi. portions of glucose, fructose, maitose, lactose, and sucrose solutions in each of five test tubes, add 5 ml of Barfoed's solution to each tube, mix, and place in a boiling water bath. Note the time when signs of reduction first appear in each tube. Continue boiling for 15 minutes, remove the tubes from the bath, and note the amounts of precipitate in the bottom of each tube after it has been standing for 15 minutes. Record your observations

Barfoed's test is not a specific test for glucose, serving simply to detect monosaccharides. Disaccharides will also respond to the test under proper conditions of acidity. Also if the sugar solution is boiled in contact with the reagent long enough to hydrolyze the disaccharide through the action of the acetic acid present in the Barfoed's solution, a positive test results. Barfoed's is a copper reduction test, but differs from Fehling's and other reduction tests in that'the reduction is brought about in an acid solution. It is unsuited for the detection of sugar in uring or in any fluid containing chlorides.

- e. Tauber and Kleiner Modification of Barfoed's Test. Introduce into one test tube 1 ml. of an approximately 0.1 per cent solution of the sugar to be tested. Put into another tube 1 ml. of water Add 1 ml. of copper reagent' to each. Heat in a boiling water bath for 3 minutes, cool for 2 minutes, Add 1 ml. of color reagent's to each Mix. A blue color will be obtained if monosaccbarides are present. With only disaccharides present the color will be the same as in the control. Chlorides interfere but not in amounts as large as 5 mg. per ml. of solution.
- f. Picric-Acid Test. To 5 ml. of the sugar solution add 2 to 3 ml. of saturated picric acid solution and about 1 ml of 10 per cent Na₂CO₂. Warm. Note the development of a mahogany-red color in the presence of glucose due to reduction of the picric acid with the formation of picramic acid:

C_4H_2 OH(NO₂)₃ \rightarrow C_6H_2 OH NH₂(NO₂)₂ Pierre seid Pieramic acid

6. Alcoholic Fermentation Prepare 500 ml of a concentrated (10 per cent) solution of glucose, add a small amount of egg albumin or commercial peptone, and introduce the mixture into a liter flask. Add yeast, and by means of a bent tube connect this flask with a second flask containing a solution of barium hydroxide protected from the air by a soda-lime tube in the stopper (Fig. 13). Place the flasks in a warm place and note the passage of gas bubbles (CO); enter, a white precipitate of barium carbonate will form. The glucose has been fermented

⁷ See Appendi

^{*}Benedict s or I olin s phosphomolybdic reagents for sugar in blood may be used See blood-sugar methods, Chapter 23

OH
$$Cu \longrightarrow CuO + H_2O$$
OH

Cupric oxide Cupric hydroxide (black) (blue)

Reaction in absence of a reducing agent

Cuprous oxide Cupric hydroxide (vellow to red)

Reaction in presence of a reducing agent

The use of a suspension of a metallic oxide or hydroxide as a reagent is obviously impractical. Certain organic compounds, particularly those containing one or more alcoholic OH groups in the molecule (e.g., tartane acid, citric acid, glycerol, even the sugars themselves), react in alkaline solution with metallic hydroxides to form a soluble complex ion which, though relatively little ionized, nevertheless dissociates to yield sufficient ions of the metal for reduction reactions to occur. The formation of such a complex is the basis of most of the heavy-metal reagents for reducing Sugars.

The alkali in these reagents hrings ahout considerable decomposition of the sugar molecule into reactive fragments which may also reduce the metal ions. Thus while the total reduction for a given concentration of sugar may be constant under carefully defined conditions and is therefore utilizable for quantitative purposes, it is impossible to write a halanced equation for the reaction in terms of the simple oxidation of the sugar and

The chemical reactions here discussed are exemplified in the following reduction of the metal ion. tests. (For the application of these and other tests to the detection of sugar in urine, see Chapter 29.)

a. Fehling's Test. To about 1 ml. of Fehling's solution in a test tube add about 4 ml. of water, and boil. This is done to determine whether the solution will of itself cause the formation of a precipitate of brownish-red cuprous the li such a precipitate forms, the Fehling's solution must not be used. To the warm Fehling's solution add sugar solution a few drops at a time and heat the mixture after each addition. The production of yellow or brownishred cuprous oxide indicates that reduction has taken place. The differences in color of the cuprous oxide precipitates under different conditions are apparently due to differences in the size of the particles, the more finely divided precipitates having a yellow color while the coarser ones are red. in the Presence of protective colloidal substances the yellow precipitate is usually formed.

in which

 $\rho = \text{sodium light.}$

 α = observed rotation in degrees,

p = g of substance dissolved in 1 ml of liquid,

l = length of the tube in decimeters

If the specific rotation has been determined and it is desired to ascertain the percentage of the substance in solution, this may be obtained by the use of the following formula

$$p = \frac{\alpha}{(\alpha)_{\nu}l}$$

The value of p multiplied by 100 will be the percentage of the substance in solution

SIECIFIC ROTATIONS OF MORE COMMON CARBOHYDRATES!

D-Glucose	+ 52 5° Sucrose	+ 66 5°
n-bructose	- 92 3° Lactore	+ 52 5°
D-Galactose	+ 81 5° Valtose	+137 0°
D-Mannose	+ 14 2° Raffinose	+101 0
L- \rabmose	+104 5° Dextrin	+19a 0°
D-Xylose	+ 19 0° Starch (soluble)	+196 0°
Rhamnose	+ 9 0° Glycogen	+197 0°
	,	±121 0

An instrument by means of which the extent of the rotation may be determined is called a polariseope or polarimeter. Such an instrument designed especially for the examination of sugar solutions is termed a saccharimeter or polarizing saccharimeter. The form of polariscope in Fig. 17 consists essentially of a long barrel provided with a Nicol prism at other end (Fig. 18). The solution under examination is contained in a tube which is placed between these two prisms. At the front end of the instrument is an adjusting eyepice for focusing and a large recording disk which registers in degrees and fractions of a degree. The light is adtirough a Nicol prism. This polarized ray that traverses the column of liquid within the tube mentioned above and if the substance is optically active the plane of the polarized ray is rotated to the right or left. Compounds rotating the ray to the right are called districted the right or left.

Sugar solutions (glucose, levulose, lactose, and maltose, but not sucrose) when freshly prepared possess a changing rotation—so-called mutarotation. For this reason such solutions before polariscopic examination should be allowed to stand over night, heated to 100° C. and then cooled, or

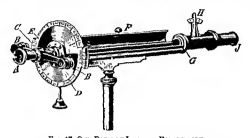


Fig. 17. One Form of Laurent Polariscope.

(B) Microscope for reading the scale; (C) a verner, (E) position of the analyzing Nicol prism, (H) polarizing Nicol prism in the tube below this point.

treated with a drop of ammonia followed by a drop of acid. This brings about an equilibrium between the α and β forms possessing different rotations, such as those of which ordinary glucose is a mixture.

Polarizing saccharimeters are also constructed by which the percentage of sugar in solution is determined by making an observation and multiplying the value of each division on a horizontal sliding scale by the value of

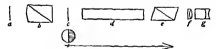


Fig. 18. Diagrammatic Representation of Course of Light through Lagrent Polariscope

The direction is reversed from that of Fig. 17. (a) Bichromate plate to purify light; (b) pointzing Nicol prism; (c) a thin quartz plate covering one-half the field and essential in producing a second polarized plane, (d) tube to contain liquid under examination; (c) analyzing Nicol prism, (f) and (g) ocular lenses,

the division expressed in terms of dextrose. This factor may vary according to the instrument.

Methods embracing the determination of the optical rotation are utilized m many analytical procedures for identifying liquids or solutions, as well as for establishing the composition of solutions.

according to the following equation

The important intermediate products formed during this reaction are indicated below (For further details see Chapter 33 on carbohy drate metabolism)

When the activity of the yeast has practically ceased, decant the supernatant fluid return it to the cleaned flask connect with a condenser and distil Catch the first portion of the distillate which may be redistilled if its alcohol content is low, and test for alcohol to the first portion of the distillate which may be redistilled in the first portion of the distillate which whi





FIG 13 FEBRENTATION AP-



FIG 14. IODOPORM

the filtrate add several drops (enough to give a yellow color to the whole mixture), of a strong solution of iodine in potassium iodide (e.g., Lugol's) Warm gently Note the iodoform odor and examine under a

gently Note the iodoform odor and examine under a microscope for crystals of iodoform. What does a positive test here indicate?

8. Demonstration of Optical Activity A demonstration I the use of the polariscope should be made by the astructor, each student being required to take readings and apply the polariscope to the determination of either the specific rotation or the concentration of a sugar.

THE POLARISCOPE

For a detailed description of the different forms of polariseopes, the method of manipulation, and the principles involved, the student is referred to any standard textbook of physics. A brief description follows

Waves of ordinary light vibrate in all planes perpendicular to the direction of propagation By certain



FIG 15 EINHORN SACCHAROMETER

means light may be caused to vibrate in but a single plane, and is then said to be plane polarized. Thus if a ray of light is passed through a crystal of calcide (natural crystallized calcium carbonate), it is divided into two rays (each polarized) vibrating in planes perpendicular to each other (Fig. 16). As the two rays are unequally hent it is possible to completely separate them and thus obtain light vibrating in but a single plane. For this purpose we may use a Nicol prism consisting of two pieces of calcide cemented together with Canada balsam. The ray of light ton entering the prism is divided into two rays. One, called the ordinary ray o, is



Fig. 16 Path of Light Ray through Nicol Prism

reflected from the Canada balsam to the exterior and absorbed by the black variish coating the prism. The other, the extraordinary ray e, passes through the balsam and emerges from the prism as polarized light. Many orgame substances (sugar, proteins, etc.) have the power of twisting, or rotating, this plane of polarized light, the extent to which the plane is rotated depending upon the nature of the molecule and upon the number of molecules which the polarized light passes. Substances which possess this power are said to be optically active. The specific rotation of a substance is the rotation expressed in degrees which is afforded by 1 g of substance tasholded in 1 ml of water in a tube 1 decimeter in length. The specific rotation, (a) b, may be calculated by means of the following formula

$$(\alpha)_p = \frac{\alpha}{p \, l}$$

As already stated, fructose, sometimes called levulose or fruit sugar, occurs widely disseminated throughout the plant kingdom in company with glucose. It has been prepared commercially from the nulin of the Jerusalem artichoke. It is not found in detectable amounts in animal tissues or fluids except under certain exceptional conditions, but in the form of its phosphoric acid esters it is a recognized intermediate in carbohydrate metabolism.

Mthough fructose is a ketose it nevertheless reduces metallic oxides in alkaline solution, owing to the presence in it of the terminal group CO CH₂OH). OF for the same reason monohydroxyacetone (CH₂ CO CH₂OH) also reduces such solutions although acetone (CH₂ CO CH₂) does not With phenylhydrazine, fructose forms the same osazone as glucose With methylphenylhydrazine, it forms an osazone more rapidly than does glucose

Fructose in solution undergoes mutarotation just as glucose does, and for similar reasons, the rotation of a fresh solution being -133 5° which changes to -92 3° on standing or in the presence of a trace of alkali. The relation between the free ketone form and the α and β forms, and the existence of pyranose and furances structures are similar to the lituation already described for glucose and are illustrated by the formulas which follow

Ordinary fructose consists chiefly of the pyranose form, fructose in the combined state as in sucrose and mulin appears to be invariably in the more reactive furanose form

EXPERIMENTS ON FRUCTOSE

I-6. Repeat Solubility, Benedict s, Phenylhydrazine, Barfoed s, Nylander s, and Fermentation tests as given for Glucose

7. Resorcinol-Hydrochloric Acid Reaction (Selivanoff). To 5 ml. of Selivanoff's reagent¹¹ in a test tube add 5 drops of a fructose solution and heat the mixture to boiling. A positive reaction is indicated by the production of a red color with or without the separation of a brown-red precipitate. The latter may be dissolved in alcohol, to which it will impart a striking red color.

To compare the reactions of aldose and ketose sugars, place 0.5-ml. portions of glucose, fructose, maltose, lactose, and sucrose solutions into each of five test tubes, add 5 ml. of Sellvanoff's reagent to each tube, mix, and place in a boiling water bath. Note the time when color first appears in each tube. Continue boiling for 15 minutes, noting the color developed in each tube at approximately 5-minute intervals. Record your observations.

This test is also given by sucrose which is hydrolyzed during the course of the test yielding fructose as one product. If the boiling be prolonged a similar reaction may be obtained with solutions of glucose or maltose. This has been explained in the case of glucose as due to the transformation of the glucose into fructose by the catalytic action of the hydrochloric acid. The precautions necessary for a positive test for levulose are as follows: The concentration of the hydrochloric acid must not be more than 12 per cent. The reaction (red color) and the precipitate must be observed after not more than 20 to 30 seconds' boiling. Glucose must not be present in amounts exceeding 2 per cent. The precipitate must be soluble in alcohol with a bright red color.

8. Aminoguanidine Reaction (Tauber). Place 0.5 ml. of concentrated sulfuric acid in a test tube. Add 0.2 ml. of a 2.5 per cent aqueous aminoguanidine sulfate solution without mixing. Add 0.2 ml. of the test solution and mix well. In the presence of ketohexoses or compounds which yield ketohexose, a bright reddish-purple color is formed in about 1 minute and persists for several hours.

As little as 0.05 mg. of fructose is said to be detectable by this test. Sucrose and inulin also give a positive test Aldohevose up to a concentration of 1 per cent does not interfere. Higher concentrations give a positive reaction, so solutions to be tested should be adjusted by dilution to contain not more than 0.5 – 1 0 per cent total carbohydrate. No color is given by aldopentoses, starch, glycogen, various pure proteins, or formaldehyde.

9. Formation of Methylphenylfructosazone. To a solution of 1.8 g. of levulose in 10 ml. of water add 4 g. of methylphenylhydrazine and enough alcohol to clarify the solution. Introduce 4 ml. of 50 per cent acetic acid and beat the mixture for 5 to 10 mlnutes (no longer) on a boiling water bath. On standing is minutes at room temperature, crystaffization begins and is complete in two hours. By scratching the sides of the flask or by inoculation, the solution quickly congeals to form a thick paste of reddish-yellow sliky needles. These are the crystals of methylphenylfructosazone. They may be recrystallized from hot 95 per cent alcohol (m. p. 153° C.). Glucose may glio the same osazone more slowly (after five hours).

¹² See Appendix,

¹³ Tauber: J. Biol. Chem., 187, 605 (1950).

GALACTOSE

Galactose occurs with glucose as one of the products of the hydrolysis of lactose. It is also found as a constituent of the galactolipides of nervous



FIG 19 MUCIC ACID CRISTALS From a 11 oto n er graph formuled by Prof. Wallass II Welker

tissue Galactose is a typical aldohexose, is dextrorotatory and exhibits mutarotation in solution and forms a characteristic osazone with phenylhydrazine It ferments either very slowly or not at all with ordinsry yeast, although some varieties of yeast ferment it resd ily Upon oxidation with nitre acid, galactose yields mucic acid (Fig 19), thus differentiating this monosacchande from glucose and fructose Lactose also yields mucic acid under these conditions Mucie acid is COOH(CHOH), COOH, the H and OH groups on carbon atoms 2 to 5 having the same spatial configuration as for galac-

tose itself Although mucic acid one half of the molecule is the mirror image of the other half (so-called internal compensation) bolling water bath for 1½ hours. Let stand aver night. A crystalline precipitate of mucic acid, which may be identified by microscopical examination of the crystals, forms under these conductions.

3. Phenylhydrazine Reaction. Make the test according to directions given for Glucose (p 63)



Mannose is an aldohevose, differing structurally from glucose solely in the spatial arrangement of H and OH groups around carbon atom number 2 (see structure, p. 58). It is found in nature chiefly in the form of polysaccharides called mannans or mannosans which occur in plants, particularly in certain seeds ("vegetable ivory"), and from which mannose may be obtained by hydrolysis. Interest in mannose has been heightened by the discovery that it is present in small amount in certain animal proteins ¹³ Mannose is a reducing sugar and is fermentable by yeast. It is readily distinguished from glucose by the formation of a sparingly soluble, colorless, crystalline phenylhydrazone when treated with phenylhydrazine at room temperature.

EXPERIMENTS ON MANNOSE

1-4 Repeat Benedict's, Barfoed's, Sellvanoff's, and Fermentation tests as given for Glucose and Fructose (pp. 62, 72)

5 Reaction with Phenylhydrazine Place a small amount of solid phenylhydrazine reagent in a test tube and add 5 ml of mannose solution Shake well and allow to stand at room temperature for 10 or 15 minutes Observe the development of a colorless crystalline precipitate of mannose phenylhydrazone (examine a drop under the microscope) When the hydrazone has been obtained, place the tube in bolimg water for one-half to three-quarters of an hour, remove, and allow to cool slowly What change has occurred? Examine a drop of the suspension at this point under the microscope Explain As a control, a glucose solution may be carried through the same experimental procedure

PENTOSES, C.H10O.

Pentoses are usually defined as sugars containing five carbon atoms in the molecule, although rhamnose, C_0H_1 O_2 , a methylpentose, is an exception to this statement. The pentoses are widely distributed in plant and anim it tissues, usually as components of some larger molecule. In plants, and more particularly in certain gums, pentoses occur as complex polymers.

Carlsberg 19 No 12 (1933)

versity of Illinois College of Medicine Smaller quantities of galactose (or lactose) may be used (40 mg) but in such cases crystallarino may take two or three days in Illinuington Biocken J 23, 430 (1929) Spiensen and Illangard Compt rend trar lab

sacchandes called pentosans, from which the free pentose (e g , arabinose, xylose) may be obtained on acid hydrolysis. In both plant and animal tissues certain pentoses (inbose, deoxynbose) are inniversally found as constituents of the nucleoproteins of the cell, being present in the nucleor acid portion of the molecule Rithose is likewise an essential component of certain mono- and dinucleotides found in cells, such as adenyhe acid, coenzymes I and II, and riboflavin (vitamin B₂)

As a class the pentoses may be either aldoses or ketoes, are nonfer mentable by yeast, have strong reducing power, and form osazones with plenythy drazine. The stereochemistry of the pentoses is similar to that already described for the hexoes. The pyranose ring appears to be the most common for the free sugars. In nucleic acids, ribose is present in the furance form. For structures of the various aldopentoses, see p. 38. On distillation with strong hydrochlone acid, pentoses and pentosans yield furfural, a reaction which is used not only for the quantitative determination of pentoses but also in the commercial production of furfural from plant by products such as out hulls.

Peutoses are an important constituent of the diet of herbivorous aminals. Their role in human nutrition is not well established. In the rare and apparently harmless condition known as pentosuma (p. 844) significant amounts of the pentose xyloketose are found in the urine and may lead to a false diagnous of diabetes mellitus. Pentosuma is also said to occur in normal individuals after the ingestion of large amounts of certain fruits.

The following experiments on pentoses may be carried out on L(+)arabinose as a typical aldopentose. The L-arabinose may be obtained from
gum arabic of from plum or cherry gum by boiling for 10 minutes with
concentrated hydrochloric acid.

just described may be considered to be derivatives of these sugars of the type known generally as glycosides. If the sugar is glucose, the compound is known specifically as a glucoside, if galactose, a galactoside, and so on The dr., tr., and polysaccharides are glycosides, as well as a wide variety of naturally occurring substances such as phlorizm, sahem, amygdalm, digitalm, etc., which contain in addition to a sugar residue a specific non-sugar portion which is known as the aglycone

Glycoside formation may be illustrated by the reaction between glucose and methyl alcohol, in which under the proper conditions a molecule of water is split off between an OH group in the glucose molecule and the

OH of the methyl alcohol, to give a methyl glucoside

Chap 2

Since the reaction involves the OH group on carbon number 1 of the glucose molecule, two methyl glucosides are possible as shown, corresponding to the α and β forms of the glucose molecule, and in general both α and β glycosides are found in nature. They may be readily distinguished by the use of enzymes which eatalyze the hydrolytic splitting of either the α or β glycoside linkage specifically. For example, in the presence of the enzyme maltase, only α glycosides are hydrolyzed, whereas the enzyme emulsin behaves similarly with respect to β glycosides. In the presence of acids both types of glycosides are readily hydrolyzed to yield their component molecules.

DISACCHARIDES

The disaccharides may be regarded as glycosides in which both components of the molecule are sugars. The common disaccharides have the general formula C₁-H₂-O₁₁, and yield hexoses on hydrolysis, a molecule of water being taken up in the reaction

$$C_{12}H_{22}O_{11} + H \cdot O = 2C_6H_{12}O_6$$

The products of hydrolysis of the more common disaccharides are as follows

Maltose = glucose + glucose Lactose = glucose + galactose Sucrose = glucose + fructose

In the formation of a gly coside linkage between two hexoses, the reducng property of one hexose is ordinarily lost. If the reducing property of the second hexose is not involved the resulting disaccharide is a reducing sugar and exhibits the general properties of such a substance (e g , osazone formation, mutarotation, etc.) Malto-e and lactose are examples of reducing disacchardes. In sucrose, however, the gly-coside inkage involves the potential reducing group of both the gluco-e and fructose components. Sucrose therefore is not a reducing sugar, does not form osazones, and does not show mutarotation in solution.

MALIOSE, CHILLION

Maltose or malt sugar is formed in the hydrolysis of starch through the action of an crzyme, vegetable amylase (diastase), contained in sprouting barley or malt Certain enzymes in the salisa and in the pancreatic jude cause a similar hydrolysis. Maltose is also an intermediate product of the action of dilute mineral acids upon starch. It is strongly dextroratory, shows mutarotation, reduces metable ions in alkaline solution, and is fermentable by yeast after being converted to glucose by the enzyme maltise of the yeast. In common with the other disaccharides maltose may be hydrolyzed by dilute acid with the formation of two molecules of monosaccharide. In this instance the products are two molecules of glucose. With phenythydrazine maltose forms an osazone, maltosazone. The following formula represents the probable structure of maltose.

D glucose 4-(a D glucoside)

As its chemical name indicates, maltose is a glucose α glucoside with a 1,4 linkage. Other disaccharides yielding only glucose on hydrolysis are known, differing from maltose in the type and position of the gluco ide bond. Thus cellobose, a glucose β -glucoside with a 1,4 linkage, is formed during the partial hydrolysis of cellulose. Gittiobiose, a rare disaccharide obtained from the roots of Gentana lutes, is a glucose β -glucoside with a 1 6 linkage. Frehalose obtained from yeast, is a nonreducing glucose a glucoside with a 11 linkage. Isomaliose, found as a minor and product of the action of amylases upon starch, is a glucose α -glucoside with a

EXPERIMENTS ON MALTOSE

1-6. Repeat Solubility. Benedict s, Nylander s, Phenylhydrazine, E # foed s, and Fermentation tests as given for Glucose, pp 62 ff

LACTOSE, C12H22O11

Lactose or milk sugar occurs ordinarily only in milk, but it has often been found in the urine of women during the period of lactation

Lactose is a reducing disaccharide, is devitor otatory, exhibits mutarotation in solution, and forms an osazone with phenylhydrazine. On hydrolysis it yields glucose and galactose. Chemical evidence indicates that the glycoside linkage involves carbon number 1 of the galactose molecule, lactose is therefore a galactoside. Enzymatic studies indicate that the galactoside linkage has the β configuration. The structure of lactose is probably as follows.

Lactose (a form)
D glucose 4 (B D-galactoside)

Since lictore exhibits mutarotation, it exists in α and β forms. The α form is the commonly obtained variety although the β form has become commercially available. It is more soluble than the α form and has been recommended for infant feeding.

I actore is not fermentable by ordinary bikers' yeast. Thus when glucore and lactose, are present together in solution they may be differentiated in terms of the reducing power of the solution before and after fermentation. On oxidation with intrie acid lactose, yields the sparingly soluble much and because of the presence of galactose in the molecule. This reaction may be used to identify lactose under the proper conditions.

In the souring of milk the I actobacillus acidophilus or Streptococcus lacticus and cert in other intercorganisms bring about lactic and ferment ation by transforming the lactoe of the milk into lactic and Cllg-CllOll CoOll this same reaction may occur in the almost try our distinct result of the action of L acidophilus and extrain other organisms. In the preparation of keft and kommiss the lactose of the milk undergoes decoholic fermentation, through the action of ferments other than yeast, and at the same time lactic and is produced.

EXPERIMENTS ON LACTOSE

Sucnose, C12H21O11

Sucrose, also called saccharose or cane sugar, is one of the most important of the sugars and occurs very extensively distributed in plants, particularly in the sugar cane, sugar beet, sugar millet, and certain palms and maples.

Sucrose is dextrorotatory, and, as before mentioned, upon hydrolysis the molecule of sucrose takes on a molecule of water and hreaks down into two molecules of monosaccharide. The monosaccharides formed in this instance are glucose and fructose. This is the reaction:

$$\begin{array}{c} C_{12} II_{22} O_{11} + II_{2} O \rightarrow C_{6} II_{12} O_{6} + C_{6} II_{12} O_{6} \\ Sucrose & Glucose & Fructose \end{array}$$

This hydrolysis may be produced by bacteria, enzymes, and certain weak acids. After hydrolysis the previously strongly dextrorotatory solution becomes levorotatory. This is due to the fact that the fructose molecule is more strongly levorotatory than the glucose molecule is dextrorotatory. The reaction is therefore frequently called inversion, and the mixture

of glucose and fructose obtained is called invert sugar.

Sucrose does not reduce metallic ions in alkaline solution and forms no osazone with phenylhydrazine. Prolonged boiling in the presence of an acid phenylhydrazine solution will, however, hydrolyzo the sucrose and cause the formation of glucosazone. It is not fermentable directly by yeast, but must first be inverted by the enzyme sucrase (invortase or invertin) contained in the yeast. The probable structure of sucrose may by represented by the following formula (Note the absence of any potential ketone or aldehyde group.) Upon inversion the active form of fructose indicated in the formula is rapidly transformed to the more stable modification.

α-D-glucopyranosyl-β-D-fructofuranoside

neutral with saturated barium hydroxide Filter off the precipitate of barium sulfate and upon the resulting fluid repeat the Phenylhydrazine, Benedict's, Nylander's, and Barfoed's reactions as given for Glucose, pp 62 ff.

TRISACCHARIDES, C18H22O16

RAFFINOSE

This trisaccharide, also called melitose or melitriose, occurs in cotton seed, Australian manna, and in the molasses from the preparation of beet sugar. It is devitrorotatory, does not reduce Benedict's solution, and is only partly fermentable by yeast.

Raffinose may be hydrolyzed by weak acids just as the polysaccharides are hydrolyzed, the products being fructose and melibiose Further hydrolysis of the melibiose yields glucose and galactose Raffinose may also be hydrolyzed by the enzyme raffinase, occurring in certain bacteria and yeasts

POLYSACCHARIDES

The polysaccharides are complex carbohydrates of high molecular weight, either quite insoluble in water or, when soluble, forming colloidal solutions Polysaccharides in the solid state do not ordinarily appear to be crystalline, but a few crystalline polysaccharides have been isolated and v-ray analysis indicates that certain polysaccharides (e.g., cellulose) possess a definite crystalline structure. Through the action of certain enzymes or of acids the polysaccharides may be hydrolyzed with the formation of simpler compounds which are regarded as constituent units of the polysaccharide. Some polysaccharides yield only simple sugars on hydrolysis, others yield not only sugars but various sugar derivatives such as glucuronic or galacturonic acid (known generally as the wrone acids), hexosamines, and even nonsugar compounds such as ethyl alcohol, sulfure acid, etc

The constituent units of the polysaceharide molecule appear to be arranged in the form of a long chain, either unbranched (cellulose, amylose) or branched (glycogen, amylopectin). The linkage between units is generally the 1,1 or 1,6 glycoside bond already described, with either the α or β configuration as the case may be Other types of linkage are known, however As a class the polysaceharides are nonfermentable and are non-reducing except for a trace of reducing power due presumably to the free reducing group at the end of a chain. They are optically active but do not exhibit mintarotation, and are relatively stable to alkali. This latter fact is utilized for example, in the separation of glycogen from tissues prior to analytical determination.

STARCH, (Call 10Os).

Starch is widely distributed throughout the vegetable kingdom, occurring in grains, fruits, and tubers. It is found in cells in the form of grainles, the micro-copic d appearance being typical for each individual starch (see p. S5). The grainites differ in size according to the source and they also differ somewhat in composition. The chief constituents are known as

amylose and amylopectin which usually exist in a proportion of about 1 3 in the granule, although some variation in this ratio may be found Amylopectin appears to contain a small amount of phosphone acid as a part of the molecule. An amylohemicellulose containing silica has also been reported as present in cereal starchus.

When boiled with water, starches form pastes. The starch granules may merely swell without disintegration and thus give a high viscosity to the solutions. Potato-starch granules disintegrate more easily and form less viscous solutions. If starch is ground in a ball mill, much of it disperseriadily in water like soluble starch. Soluble starch is formed by the action

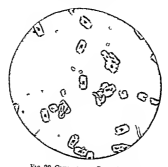


Fig 20 Chystalline Corn haylose, (Kerr)

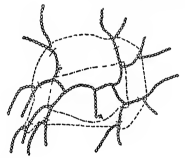
of dilute hydrochlone and upon starch. It is so called because it readily forms a imput clear solution with hot water. The and apparently modifies the amylopectin of the granules so that it disintegrates more completely on heating with water.

On hydrolysis with acids starch yields glucose. The two major constituents of the starch granule amylose and amylopectin, must differ largely, therefore in their molecular structure. The amylose molecule appears to consist of about 300 glucose units in an unbranched chain, with a molecular weight of about 30 000. When pure amylose is sparingly soluble in water and forms a clear solution. It gives a deep blue color with iodine, and is completely hydrolyzed to maltose by the enzyme β -amylase. Amylose has been obtained in the crystalline form by Kerr (Fig. 20). Amylopectin on the other hand forms opalescent volutions, gives a purple to violet color with iodine which is much less intense than in the amylose reaction, and is only partly hydrolyzed to maltose by β -amylase. Amylopectin appears to contain many hranched glucose chains in the molecule.

each chain having about 25 glucose units, and the entire molecule having a molecular weight up to 20 times that of amylose (Fig. 21).

The linkage between glucose units in the unbranched amylose chain is the 1,4 glycoside bond described previously. The structure of a portion of such a chain is probably that indicated in the formula:

In the branched-chain structure characteristic of amylopectin, both 1,4 and 1,6 glycoside bonds are found; the 1,4 linkage forms the linear por-



A Aldah die and

(

A Aldehydie end-group.

 End of initial degradation by β-any lase, yielding residual-dextrix I.

Limit of degradation produced by 2-glacosidase, giving dextrin II, hydrolyzable by β-amylase.

oy φ-anylase.

End of further attack by β-amylase, yielding residual dextrin III.

Fig. 21. Meyen's Schematic Representation of the Brancheo Amylopectin Molecule (Handle), Courters, Wallerstein Communications.

tion of a chain and branching takes place by mion of a side chain through carbon atom number 1 of a terminal glucose residue of the side chain with carbon atom number 6 of a glucose residue in the main chain.

In the course of the digestion of starch by salivary or pancreatic α -amplase there is first formed soluble starch (a clear solution giving a blue

color with iodine) then dextrins giving blue or red colors with iodine, next achrodextrins giving no color with iodine, and finally maltose. Some maltose is, however, formed almost from the beginning of the digestion. The amylase apparently catalyzes the hydrolytic splitting of every other glycoside hond thus producing maltose units. The action of a amylase on the linear and branched glucose claims, amylose and amylopeetin, respectively, is discussed on p. 352. In the case of acid hydrolysis the same intermediate products are formed but gluco e is the end product. The hydrolysis of starch by acid is an example of the catalytic action of the hydrolysis of starch by acid is an example of the catalytic action of the

Synthetic starches have been prepared by the action of certain muscle and potato enzymes on clucose 1 phosphate, the Cornester (see p. 333 for an experiment demonstrating such a synthesis). These synthetic starches results are the results are the synthesis.



Fig 22 Potato



FIG 23 BEAN



FIG 24 ARROWNOOT



Fig 25 Rye.



FIG 26 BARLEY



Fig 27 OAT



Fic 28 BUCKWIEAT



Fig 29 MAIZE



Fig 30 Rick



Fig. 31 Fey. The 32 Wheat Fig. 22/32 Struck Chancles from Various Sources (beforen by Fey.)



- 6. Benedict's Test On starch solution²⁷ (see p. 66). Does starch have any detectable reducing ability?
- 7. Hydrolysis of Starch. Place about 25 ml. of 1 per cent starch solution II na small beaker, add 10 drops of concentrated HCI, and boil gently. By means of a small pipet, at the end of each minute remove a drop of the solution to the test tablet and make the regular lodine test. At the end of the same 1-minute intervals add exactly 3 drops of the mixture to 5-ml. portions of Benedic's solution in a series of test tubes As the testing proceeds the reaction with lodine should become weaker and finally be negative. At this point place all the tubes containing Benedict's solution in a boiling water bath for 3 minutes, then remove and allow to cool Note the degree of reduction in each case and compare with the rate of disappearance of the iodine reaction. Make the phenyihydrazine test upon some of the hydrolyzed starch. What sugar bas been formed?
 - Diffusibility of Starch Paste. Test the diffusibility of starch through a suitable dialyzing membrane. Compare with glucose in this respect.

INULIN, (CallioOs),

Inulin is a polysacchande which may be obtained as a white, odorless, tasteless powder from the tubers of the artichoke, elecampane, or dablia It has also been prepared from the roots of chicory, dandelion, and burdock. The rubber-producing plant guayule also contains nulin. It is very slightly soluble in cold water and quite easily soluble in hot water. In cold alcohol of 60 per cent or over it is practically insoluble. Inulin gives a negative reaction with iodine solution. It is very difficult to prepare inulin which does not reduce Benedict's solution slightly. This reducing power may be due to an impurity Practically all commercial preparations of inulin possess considerable reducing power Inulin is a polymenzed form of fructofurances, containing about 30 fructofurances units per chain, united by 1,2 linkages.

laulin is levorotatory and upon hydrolysis by acids or by the enzyme inulase it yields the monosacchande fructose which readily reduces Benedict's solution. The preparation of fructose by hydrolysis of the inulin of the Jerusalem artichoke has become of commercial importance. The ordinary amylolyte enzymes occurring in the animal body do not digest inulin. A small part of the ingested mulin may be hydrolyzed by the acid gastine junce, but the value of nutin as a significant source of energy in human dietanes must be questioned. Bats may form some glycogen from nulin-lindin administered intravenously is readily excreted by the kidneys, aptopartity because the mulin molecule though colloidal, is sufficiently small this property in the mulin delarance test of kidney function and in other studies of renal physiology.

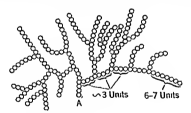
 Iodine Test. (a) Place 2 to 3 ml. of the inulin solution in a test tube and add a drop of dilute iodine solution. Compare with a control containing water instead of inulin solution. What do you observe?

(b) Place a small amount of inulin powder in one of the depressions of a test tablet and add a drop of dilute lodine solution. Is the effect any different from that observed above?

- 3. Resorcinol-Hydrochloric Acid Test. Test a portion of inulin solution by this test, following the directions given on p. 73. Explain the results.
- 4. Benedict's Test. Make this test on the inulin solution according to the instructions given for Glucose (p. 66). Is there any reduction? Explain.
- 5. Hydrolysis of Inulin. Place 5 ml. of inulin solution in a test tube, add a drop of concentrated hydrochloric acid, and boil for one minute. Now cool the solution, neutralize it with concentrated sodium carbonate solution, and test the reducing action of i ml. of the solution upon I ml. of Benedict's solution. Compare with a control using I ml. of unbydrolyzed inulin solution. Also try the Resorcinol-Hydrochloric Acid reaction as given on p. 73, likewise comparing with a control of untreated inulin solution. Explain the results.

GLYCOGEN, (C6H10O5)x

Glycogen is the form in which carbohydrate is stored in the animal organism. It is found in the liver, muscles, kidneys, and other ussues, but is



F10 33 STRUCTURE OF GLYCOGEN.

The circles represent glucose units, (A) the aldeby die end of a chain.

Meyer. Advances in Britingloon, 3, 199, 1943

notably absent from brain. On hydrolysis with amylase it yields maltose, and with acid it yields glucose. It thus resembles starch and the destrius in these respects but differs significantly from these compounds in molecular architecture. In the glycogen molecule the component glucose residues are linked in chains by both 1,1- and 1,6- a-glycoside bonds as with the starches, to give a highly branched structure as contrasted with the unbranched amyloso or slightly branched amylopectin chains of starch (see Fig. 33). Glycogen is soluble in cold water to form an opalescent solution and ordinarily gives a red color with iodine, although some forms of

glycogen which give blue or purple colors with induit are known. These color differences are apparently related to the extent of chain branching the blue color representing relatively unbranched chains and the red color with induite corresponding to highly branched chains.

For a further discussion of Llycogen, and experiments, see pp 272

and 280

DEXTRIN, (C₆H₁₀O₆),

The dextrins are found as intermediate products in the course of hy drolysis of starch to glucose or maltose by acids or enzymes. They are colloidal in nature but of a lower degree of molecular complexity than starch as would be expected a variety of dextrins are known, most of which are relatively ill-defined, the higher dextrins resembling starch in certain respects while the lower dextrins more nearly resemble the sugars. As a class the dextrins are readily soluble in water, insoluble in alcohol, and do not diffuse through cellulose membranes of ordinary porosity.

The dextrins may be hydrolyzed by dilute acids to form flucose and by amylases to form maltose. They are not fermentable by yeast. Various dextrins give different colors with iodine, depending apparently upon their molecular complexity the higher dextrins give a blue or purple color, in termediate dextrins give a red color, and the lower dextrins may give no color at all. As is the ease with most other polysaccharides, the glucose chains in the dextrin molecule have a free reducing group at one end the ability of dextrins to reduce Benedict's solution therefore depends upon the molecular weight and chain organization, dextrins with low molecular weight showing detectable reducing power. The reducing ability of commercial dextrin preparations is ordinarily due to the presence of free sugar. The formation of various dextrins as intermediate products of the action of amylases upon starch is illustrated schematically in Fig. 21.

bulk of saturated ammonium sulfate, shake vigorously, and allow to stand for five minutes. The starch is precipitated. Fifter through a dry paper, and to a portion of the filtrate add a drop or two of lodine solution. Compare with an iodine test on the detrin solution alone which has been treated with ammonium sulfate in like manner.

89

- 4 Benedict's Test See if the dextrin solution will reduce Benedict's solution
- 5 Precipitation by Alcohol To about 50 ml of 95 per cent alcohol in a small beaker add about 10 ml of a concentrated dextrin solution Dextrin is thrown out of solution as a gummy white precipitate
 - 6 Diffusibility of Dextrin (See Starch, 8, p 86)

insolubility of cellulose

Cellulose forms a large portion of the cell walls of plants. Physically, cellulose is distinguished from other carbohydrates largely by its extreme insolubility in most of the ordinary solvents. Chemically, cellulose has been shown to consist of a large number of β glucose units jouned together in a chain by 1,4 glycoside bonds, as indicated in the formula given herewith Various methods for estimating the number of glucose units in the cellulose

molecule yield results which vary from 200 to 2,000 such units, depending upon the method employed. A ray studies of cellulose indicate that it is crystalline in structure and consists of bundles of long chains with the chains running parallel to the fiber axis. These chains are unbranched and straight rather than coiled, and there is some evidence for the existence of cross linkages between adjacent chains the chemical nature of such cross linkages is obscure. The close packing of the long straight chains produces a fiber with much mechanical strength, and may also account for the

Cellulose is not soluble in water or the usual organic solvents nor in dilute acid or alkali. It is soluble however in a variety of special solvents, such as Schweitzer's reagent (ammoniaeal copper hydroxido), zine chloride—hydrochlonic acid solution, and sodium hydroxide and CS- which latter are the reagents for making viscose, a form of artificial silk. With 10 per cent sodium hydroxide, cellulose is converted into hydrocellulose and is said to be increarized. About 70 per cent silfuric acid converts it into regetable porchiment. Such commercial products as rayon cellophane, sausage caling, etc., are made from cellulose. With strong nitne acid and concentrated sulfuric acid it forms introcellulose.

Cellulose is not hydrolyzed by boiling with dilute mineral acids. It may

be hydrolyzed, however, by treating with concentrated sulfuric acid, their diluting the solution with water, and boiling. The product of this hydrolysis is glucose. Hydrolysis of cellulose by certain bacteria has been reported to yield the disaccharide cellulose, analogous to the production of maltose from starch.

There is some difference of opinion as to the exact extent to which cellilose is utilized in the animal organism. It is no doubt more efficiently utlized by herbivora than by carnivar or by man. It is claimed that about 25 per cent may be utilized by hierbivora and less than 5 per cent by dogs, whereas the quantity utilized by man is too small for it to play a nutrtional role in the diet of a normal individual. In neither man nor the lower animals has there been demonstrated any formation of sugar or glycoger from cellulose. It is probable that the cellulose which disappears from the intestine is transformed for the most part into lower fatty acids (acetic etc.) by the action of intestinal bacteria. A cellulase capable of digesting awdust and filter paper has been found in the digestive diverticula attached to the stomach of the shipworm.

EXPERIMENTS ON CELLULOSE

For these experiments, a high-grade filter paper, absorbent cotton, ϵ cleansing tis ue may he used

- Solubility Test the solubility of cellulose in water, dilute and concentrated acid, and alkali
- 2 Iodine Test Add a drop of dilute lodine solution to a few shreds of \cot^{co} on a test tablet Cellulose differs from starch and dextrin in giving no \cot^{co} with lodine
- 3 Formation of Amyloid ** To 6 ml of distilled water in a test tube, add 16 ml of concentrated sulfuric acid. The acid should be added to the water in small portions and the mixture stirred with a stirring rod and cooled under the tap, or by immersion in a beaker of cold water, between additions. To the cooled mixture add at two inch square of cleansing tissue and stir for from 5 to 10 minutes, when most of the tissue is dissolved. Pour about 3 ml of this solution into about 10 ml of distilled water, and note the flocculent precipitate of amyloid formed. To another small portion of the solution add lodine and note the blue or black color formed. Pour the remainder of the acid solution of tissue into about 25 ml of distilled water in a small beaker and host for 15 to 30 minutes. Now cool, neutralize with solid sodium carbonate, and test with Benedict's solution. Glucose has been formed from the cellulose by the action of the acid.

- 5. Hydrochloric Acid—Zinc Chloride Solubility Test (Cross and Bevan). Piace a little absorbent cotton in a test tube, add Cross and Bevan's reagent, is and stir the cellulose with a giass rod. When solution is complete, reprecipitate the cellulose with 95 per cent alcohol.
- 6. Iodine—Zinc Chloride Reaction. Place a little absorbent cotton or quantitative filter paper in a test tube and treat it with the iodine—zinc chloride reagent.¹⁹ A blue color forms on standing. Amyloid has been formed from the cellulose through the action of the ZnCl₂ and the iodine solution has stained the amyloid blue.
- 7. Other Cellulose Solvents. It has been demonstrated by Deming that there are many excellent solvents for cellulose (filter paper)—for example, the concentrated aqueous solutions of certain salts such as (1) antimony trichloride, (2) stannous chloride, and (3) zinc bromide. In hydrochloric acid solution the solvent action of the above saits is increased. The following salts are also good solvents in hydrochloric acid solution: mercuric chloride, bismuth chloride, antimony pentachloride, tin tetrachloride, and titanium tetrachloride. In the case of the last-mentioned salt the swollen, transparent character of the celiulose fibers preliminary to solution can be seen very nicely.

Try selected solvents suggested by the instructor.

HEMICELLULOSES

The hemicelluloses differ from cellulose in that they may be hydrolyzed upon boiling with dilute mineral acids. They differ from other polysac-charides in being not readily digested by amylases. Upon hydrolysis hemicellulose may yield pentoses or hexoses or both, together with uronic acids. The vegetable gums and pectius may be included under this head.

Pentosans. Pentosans yield pentoses upon hydrolysis. So far as is known they do not occur in the animal kingdom. They have, however, a very wide distribution in the vegetable kingdom, being present in leaves, roots, seeds, and stems of all forms of plants, many times in intimate association or even chemical combination with galactans and uronic acids. In herbivora, pentosans are 40 to 80 per cent utilized. The few tests on record as to the peutosan utilization by man indicate that 80 to 95 per cent disappears from the intestine. According to Cramer, bacteria are efficient hemicallulose transformers. It has not yet been demonstrated that pentosans form glycogen in man, and for this reason they must be considered to play an unimportant part in human nutrition. Gum arabic, an important pentosan, may be hydrolyzed by boiling with strong hydrochloric acid for a short time. The pentose arabinose results from such hydrolysis.

Galactans. In common with the pentosans the galactans have a very wide distribution in the vegetable kingdom. One of the most important members of the galactan group is agar-agur, a product prepared from certain types of Asiatic or American seaweed. Chemically, the agar molecule appears to consist of a chain of p-galactose units linked by a 1,3-glycoside bond, with a single L-galactose unit in a 1,1 linkage at the reducing end of the chain, this latter unit also being esterified in the position with sulfuric acid. Thus the products of hydrolysis include much p-galactose, some L-galactose, and sulfuric acid. This galactan is about 50

per cent utilizable by herhivora and 8 to 27 per cent utilizable by man Agar ingestion bas been shown to be a very efficient therapeutic aid is cases of chronic constipation. This is particularly true when the constipation is due to the formation of dry, hard feeal masses (scybala), a type of feeal formation which frequently follows the ingestion of a diet which is very thoroughly digested and absorbed. The agar, because of its relative indigestibility and its property of absorbing water, yields a bully feeal mass which is sufficiently soft to permit of casy evacuation. Agar has heen used with good results in the treatment of constipation in children. Agar is not limited to its use in connection with constipation, but may serve in other capacities as an aid to intestinal therapeutics.

Dextrans and Levans. The dextrans are polysaccharides produced by the action of certain bacteria on sucrose, the reaction being as follows:

$$nC_{12}H_{22}O_{11} = nC_6H_{12}O_6 + (C_6H_{10}O_5)$$

Sucrose Fructose Dextran

In effect, the fructose residue is removed from the sucrose molecule, and the remaining glucose residues polymerized to form the polysaccharide dextran. The dextrans bave a high molecular weight (which varies with the conditions of preparation and species of organism used) and form rather viscous colloidal solutions in water. For many years they were regarded as undesirable by-products in the sugar and fermentation industries. They have attracted considerable clinical interest because of their proposed used as plasma substitutes, since they form nontoxic solutions with a high colloidal osmotic pressure, suitable for intravenous administration as a temporary replacement for plasma proteins lost from the blood by hemorrhage. Dextrans so administered gradually disappear from the circulation; their metabolic fate is however still obscure.

The dextrans consist almost entirely of p-glucose units joined in chains by 1,6-glycoside bonds, with evidence for occasional cross-linking between chains by the formation of 1,4 bonds. On bydrolysis with dilute acids, dextrans yield only p-glucose. In general, dextrans are not acted upon by ordinary amylases, although some exceptions to this rule are known.

Similar polysaccharides containing fructose residues and known as levans are also produced by certain microorganisms. The reaction is analogous to that given for dextran formation, with the formation of glucose and polymerized fructose (levan), except that the incidence of side reactions involving the hreakdown of part of the sucrose to free glucose and fructose appears to he greater. Chemically, levans are found to he polymers of fructose with 2.6 linkages between units, thus differing from inulin, which contains 1.2 linkages. Levan formation has relatively little.

The Peetins. The peetins are colloidal earbohydrates which with the proper concentration of acid and of sugar form gels. For a gel to form, there must be present from 0.3 to 0.7 per cent peetin, 65 to 70 per cent of sugar (usually sucrose), and a pH of 3.2 to 3.5. Commercial peetin is prepared from apples and lemons. On hydrolysis the peetins yield galacture.

ronic acid, arabinose, galactose, acetic acid, and methyl alcohol. The characteristic properties of pectin appear related to the presence of a long chain of anhydrogalacturonide residues, partly methyl esterified. Nongalacturonide material, galactan, and araban, which appear to act chiefly as diluents, may however make up a considerable portion of the weight of the pectin.

EXPERIMENTS ON A PENTOSAN

- Solubility. Test the solubility of gum arabic in hot and cold water and alcohol.
- Iodine Test. Add a drop of dilute iodine solution to a little gum arabic on a test tablet. It resembles celluiose in giving no color with iodine.
- 3. Hydrolysis of Gum Arobic. Introduce a little gum arabic into a test tube, add 5 to 10 ml. of strong hydrochloric acid (conc. HCl and water 1:1), and heat to bolling for 5 to 10 minutes. Cool, neutralize with sodium hydroxide, and test hy the Benedict or some other reduction test. A positive reaction should be obtained, indicating that the gum arabic has been hydrolyzed by the acid with the production of a reducing substance. What is this reducing substance? How would you identify it?

EXPERIMENTS ON A GALACTAN

- Solubility. Test the solubility of agar-agar in hot and cold water. Observe its marked property of imbiling water (see above).
- 2. Iodine Test. Add a drop of dilute iodine solution to a little agar-agar on a test tablet. It resembles cellulose in giving no color with iodine.
- 3. Hydrolysis of Agor-ogor. Introduce a few pieces of agar-agar in a test tube, add 5 to 10 ml. of strong bydrochioric acid (conc. HCl and water 1:1) and heat to bolling for 5 to 10 minutes. Cool, neutralize with sodium hydroxide, and test by the Benedict or some other reduction test. A positive reaction should be obtained, indicating that the agar-agar has been hydrolyzed by the acid with the production of a reducing substance. What is this reducing substance? How would you identify it?

EXPERIMENTS ON PECTIN

- I. Preparation of Pectin. Pare off the yeilow layer from a grapefruit rind. Run through a meat chopper, cover with water, and let stand over night. Strain on cheesecloth and squeeze out the fluid. Boil the pulp with water for about two hours, hinging finally to a low volume. Pour off the fluid and add alcohol to precipitate the pectin. Filter and dry.
- 2. Formotion of a Gcl. Into a weighed 400-ml. heaker introduce 70 g. of cane sugar, 1 g. of dry pectin, 0.5 g. of citric or tarturic acid, and 100 ml. of water. Heat to boiling. Concentrate to a weight of 100 g. Let it stand over night.

REVIEW OF CARBOHYDRATES

In order to facilitate the student's review of the carbohydrates, the preparation of a chart similar to the appended model is recommended. The signs + and - may be conveniently used to indicate positive and nega-

Fats

THE LIPIDES

The term lipide is applied to a group of unturally occurring substances characterized by their insolubility in water and their solubility in such fat solvents as ether, chloroform, boiling alcohol, and benzene. They are limited to substances which are utilizable by the animal organism. Individual members of this group show large individual variations in solubility, but as a class the lipides are readily distinguishable from the carbohydrates and the proteins, the other two great groups of naturally occurring compounds. Chemically, the lipides are either esters of fatty acids or substances capable of forming such esters. They are very widespread in nature, being found in all vegetable and animal matter. Some memhers of this group, such as the phosphatides and sterols, are found in all living cells where, with the proteins and carbohydrates, they form an essential part of the colloidal complex of protoplasm. Complex lipides are also found in large quantities in hrain and nervous tissues, thus indicating the important role these substances must play in the living organism. Other lipides, such as the fats and oils, represent the chief form in which excess nutrients are stored in the animal body. They arise from ingested lipides and from the metabolism of earbohydrates and proteins. and are stored in fat deposits, such as in the subcutaneous connective tissue, in the intermuscular connective tissue, in the omentum, in the perirenal fat depots, and in the genital fat. Lipides act as heat insulators and as reserve supplies of energy. They may be classified as follows:

CLASSIFICATION

SIMPLE LIPINES

The simple lipides are esters of fatty acids with certain alcohols. They are usually further classified according to the nature of the alcohols, as follows:

1. Fats and Oils. Esters of fatty acids and glycerol. Oils are fats which

are liquid at room temperature.

 Waxes, Esters of fatty acids with long-chain aliphatic alcohols or with cyclic alcohols. These may be subdivided into: (1) True waxes; (2) Cholesterol esters; (3) Vitamin A and its carotenol esters; and (4) Vitamin D esters.

COMPOUND OR CONJUGATE LIMIDES

The compound lipides are esters of fatty acids which, on hydrolysis, yield other substances in addition to fatty acids and an alcohol Some important members of this group are

1. Phosphollpides (Phosphatides). I ipides which, on hydrolysis yield fatty acids, phosphoric acid, sometimes, but not always, glycerol, and a nitrogenous base These are subdivided into the following groups

(a) LECITHIAS Lipides containing fatty acids, phosphoric acid, glyc-

erol and the nitrogenous hase choline

(b) CFPHALINS Lapides which yield on hydrolysis fatty acids, glycerol phosphoric acid, and either the hase ethanolamine (colamine) or the amino acid serine Lipides of uncertain structure which contain mositol fatty acids, phosphoric acid, ethanolamine and possibly galactose and tartaric acid are also included in this class

(c) Sphingouyelins Lipides containing the introcenous base sphin gosine a single fatty acid molecule, phosphoric acid and choline, but no glycerol

2 Cerebrosides. Lipides which contain carbohydrate (galactose or glucose), one fatty acid and sphingosine, but no phosphoric acid or glyccrol 3 Sulfolipides Similar to cerebrosides except that sulfuric acid is

present as cerebronic acid ester Sulfolipides containing sphingosine, ga lactose cerebronic acid, sulfuric acid and potassium have been described 1

DERIVED LIPIDI 9

The derived lipides are substances formed in the hydrolysis of simple or compound lipides which still retain the properties of this class of compound

1. Fatty Acids Saturated and unsaturated acids

2 Alcohols Compounds of high molecular weight hut not gycerol These may be classified as follows

(a) ALIPHATIC ALCOHOLS such as cetyl (CH2(CH2)14CH2OH) stearyl (CH₂(CH₂)₁₆CH₂OH) and myricyl (CH₂(CH₂)₂₆CH₂OH)

(b) STEROLS which contain the phenanthrene nucleus (cholesterol ergosterol, sitosterol stigmasterol)

(c) Alcohols Containing the \$ long Ring These include vitamin A, kitol and carotenols such as cryptoxanthin lutein and zeaxanthin

3. Hydrocarbons Compounds having no earboxyl or alcohol groups, and which cannot be saponified

(a) ALHERATIC HYDROCARDONS Pentarosane CH4(CH2)22CH2, and homologues to hentriacontane Callie

(b) Carotevoids C40H44 compounds such as α, β- and γ-carotere and by copene

(c) Squaling Unsaturated hydrocarbon Caollie in olive and shark liver oils

4 Vitamins D Differ from steroids in that the phenanthrene nucleus is ruptured between carbons 9 and 10

¹ Blix Z physiol Chem 219 82 (1933)

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5. Vitamins E. Chroman derivatives, α-, β-, γ-, and δ-tocopherols 6. Vitamins K. Derivatives of 1.4 naphthogumone with long hydrocarbon side chains

THE FATS

The Fatty Acids. The fatty acids found in fats and other lipides are of various types Some of them, like palmitic acid (CH3(CH2)14COOH) and stearic acid (CH₂(CH₂)₁₆COOH), are straight-chain saturated acids lielonging to the acetic acid series, and have the general formula $C_\alpha H_{2\alpha} O_2$ Others are unsaturated and have from one to four, and even more, double bonds to their molecules. Thus olete acid $(C_{18}H_{24}O_2)$ has one double bond in its molecule, linoleie acid (C₁₈H₂·O₂) has two double bonds, and hnoleme acid (C18H30O2) has three Practically all the fatty acids found in nature contain an even number of carbon atoms, although porpose oil and the head oil of the white whale have been shown to contain isovaleric acid (C₆H₁₀O₂) In addition to the above types of acid, some hpides contain hydroxy acids both saturated and unsaturated, and dicarboxyhe acids Thus castor oil contains an unsaturated hydroxy acid, ricinoleic acid (C18H34O3) Certain cyclic saturated and unsaturated fatty acids are hkewise found in nature Some of the more important chemical and physical properties of saturated and unsaturated fatty acids are discussed below

SATURATED FATTY ACIDS, C. H2.O2 OR C. H2.+1COOH

The physical properties of the saturated fatty acids depend upon their molecular weights Whereas those fatty acids that contain ten carbon atoms or fewer in their molecules are hauds at room temperature, the remainder are solids whose melting points rise with increasing molecular weight The liquid acids are also known as volatile fatty acids, since they may be distilled with steam, whereas the others, the nonvolatile acids, are

SATURATED FATTY ACIDS

SATURATED FATTY ACIDS				
Common Name	Chemical Name	Structure	Occurrence	
Buty rie Caproie Capry lie Caprie Laurie	Butanoic Hexanoic Octanoic Decanoic Dodecanoic	CH ₄ (CH ₂) ₂ COOH CH ₂ (CH ₂) ₄ COOH CH ₄ (CH ₂) ₄ COOH CH ₄ (CH ₂) ₄ COOH CH ₄ (CH ₂) ₄ COOH	Butter fat Butter fat coconut oil Butter fat coconut oil Butter fat coconut oil Laurel kernel oil butter fat, coconut oil	
Vi3 ristic	Tetradecanoic	CII. (CII.)., COOII	Nutineg fat, butter fat	
Pulmitic	Hevadecanoic	CII. (CII.)1. COOII	Most vegetable and animal fats Vlost vegetable and animal	
Steame	Octadecanoic	CII. (CII.) 14 COOH	fats	
Vrachidie Behenie Lignocerie	I teo more Docosanore Tetracosanore	CII. (CII.) 1 COOH CII. (CII.) 2 COOH	Pennut oil Rapesced oil, peanut oil Cercbrosides sphingomyelin peanut oil	

carried over by steam distillation only in traces or not at all Fatty acids with four carbon atoms or fewer are miscible with water in all proportions As the length of the carbon chain increases beyond this, however, the solubility rapidly diminishes to zero. The common straight-chain saturated fatty acids found in nature as constituents of lipide molecules are hated in the table on p 99

UNSATURATED FATTY ACIDS

The unsaturated fatty acids are characterized by the presence of one or more double honds in the molecule. They have been classified in accordance with the number of double bonds as monoethenoid, diethenoid, triethenoid, etc, and named by reference to the parent hydrocarbon, the position of the double bond or bonds in the chain being indicated by a number referred to the carboxyl carbon atom as number one As is true of the saturated fatty acids, many of the unsaturated fatty acids have com mon names which may be used as frequently as the chemical name

Because of the presence of the double bond the unsaturated fatty acids are much more reactive than the saturated acids, the reactivity increasing with increase in the number of double honds. The unsaturated fatty acids are capable of taking up one molecule of water, oxygen, hydrogen, hromine, or iodine at each double hond, and the amount of such substance (e g , iodine) absorbed by a given weight of acid is used to determine its degree of unsaturation It is obvious that a variety of isomerism is possible among the unsaturated fatty acids, depending not only on the posi tion of the double bond in the chain but also on cis trans isomerism across a double bond Relatively few of the large number of possible isomers of the unsaturated fatty acids are found in nature

The most common unsaturated fatty acid found in nature is the monoethenoid acid olcic acid (9-octadecenoic acid) This acid is so widely distributed that, according to Hilditch, no natural fat or phosphatide has as yet been found not to contain oldie acid Vaccenic acid, an isomer of oleic acid, has received considerable attention. It occurs in low concentration in certain animal and vegetable tissues. Other common unsaturated fatty acids include (1) the diethenoid acid linoleic acid, found in both animal and plant fats, (2) linolenic acid, which is a triethenoid acid found largely in vegetable fats, but also in some animal fats (horse, egg yolk), and (3) arachidonic acid, a tetraethenoid acid which is found in both the fat and the phosphatide fractions of many animal tissues, particularly in liver and in suprarenal phospholipides. The chemical characteristics of these representative straight-chain unsaturated fatty acids tollow.

Cor mon Name	Empirical Formula	Number of Double Bonds	Chemical Vame
Oleie Veid Lincoeie Veid Lanoet ie Veid Vrachistonie Veid	C,II,O, C,II,O, C,II,O,	2	9-Octadecenoic 9-12-Octadecadienoic 9-12-15-Octades strict ic 5-8-11-14-I icosatetra noic

From the empirical formulas it can be seen that olde, hnolder, and linelenic acids may be considered derivatives of the saturated C₁₈ acid, stearic acid, and arachidonic acid is an unsaturated form of the C₂₀ acid, arachidic acid Similar ethenoid derivatives are known of many of the other saturated fatty acids found in nature

In addition to the straight-chain fatty acids so far described, a number of branched-chain and cyclic fatty acids, both saturated and unsaturated, have been isolated from natural sources. This erculostearic acid (10-methylistearic acid) has been obtained from the wax of the tubercle bacillus, as has phthioic acid, a methylated branched chain C₂₅ acid which is possibly 3,13,19-trimethyltricosanoic acid, although this structure has been disputed. The latter appears to be associated with some of the clinical manifestations of tuberculosis. It is interesting to note that these and other fatty acids are found in tubercle wax as esters of the disaccharde trehalose (see Chapter 2). In chaulmoogra oil the unsaturated cyclic fatty acids chaulmoogra acid and hydnocarpic acid are found, these acids or their derivatives have been used in the treatment of leprosy.

The Fats. The fats are neutral esters of glycerol and fatty acids. An example is tristearin, synthesized in living tissues from one molecule of glycerol and three molecules of stearic acid.

Of the three molecules of fatty and entering into the composition of a "true" or neutral fat, all may be the same, as in tristearin, in which case it is referred to as a simple triglyceride. However, more commonly two or three different fatty ands are involved, and such a fat is called a mixed glyceride, the composition being midicated by the name as oleodipalmitin, stearodiolem, oleopalmitostearin, etc. Mixed glycerides occur much more commonly than do simple glycerides such as tristearin, tripalmitin, and triolem. The possibilities of stereosomerism among the fats are many, they include isomerism due to the same fatty acids being arranged differently in the molecule, as well as optical isomerism when the middle carbon atom of the glycerol portion becomes asymmetric or when an optically active fatty acid is present. Such isomerism appears to be of much less physiological importance in the fats than in carbohydrates and proteins. Diglycerides (containing two fatty reads) and monoglycerides (containing one fatty acid) have been shown to be normal intermediates arising

in fat digestion. The 1,2-digly ceride and the 2-monogly ceride are believed

to be the primary types,2

Animal Fats. Naturally occurring animal fats consist largely of mixed glycrides of oleic, palmitte, and steare acid; furthermore they are usually mixtures of individual fats. Fats from various sources differ considerably in their fatty acid composition. Mutton fat contains more stearic acid and less oleic acid than pork fat. Human fats contain a high percentage of oleic acid. Butter fat consists largely of glycerides of palmitic and oleic acids, with small amounts of stearic acid and of the lower fatty acids such as butyne and caproic; significant variation in this connection may be noted from species to species (see Chapter 12).

General Properties of Fats. Pure neutral fats are odorless, tasteless, and generally colorless; the color of natural fats and oils is ordinarily due





hig 35 Metros FAT.

Although most fats may exhibit sharp melting points under certain specified conditions marked differences can be demonstrated between melting point and solidifying point. These discrepancies have been related to the fact that fats may occur in several polymorphic forms, depending upon the rate of cooling and the temperature at which they are maintained The several forms are referred to as α , β , β' , and γ types. They vary in stability, melting point, gross and microscopic appearance, \ ray diffraction pattern, erystal structure, density, and other properties. The y form has the lowest melting point, the α , β (when present) and β forms have progressively higher melting points. The polymorphic forms change to the B isomorph on standing, this is the most stable of the polymorphic

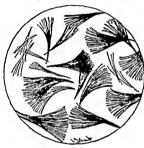


FIG 36 PORK FAT

forms Terguson and Lutton' reviewed the polymorphism of fats, and the melting points of the known modifications have been listed by Bailey 5

Mixed glycerides contuming a high proportion of unsaturated fatty acids are usually liquid at room temperature, as indicated above, and are commonly called oils These oils take up hydrogen at their double bonds, in the presence of catalysts such as finely divided nickel, and are thus converted into solid fats. This process is called hardening or hydrogenation Many commercial fats are partially hydrogenated vegetable oils There is no known nutritional objection to the use of properly hydrogenated rejetable oils in place of animal fats. For example, in experiments on rats and on man it was shown that hydrogenated vegetable oil was as satisfactorily digested and utilized as lard, and was less hable to cause gastric or intestinal symptoms of an objectionable nature 6 Furthermore, under conditions in which oxidation may cause destruction of essential factors in the diet, the use of hydrogenated oil is claimed to be preferable to the use of unhydrogenated vegetable or animal fats 7

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When a fat is allowed to stand for a sufficient length of time in contact with air and moisture particularly in the presence of light and heat cer tain changes occur and it becomes rancid Three types of rancidity have been described, namely, oxidative, ketonic and hydrolytic The first is the most important in the spoilage of fat Rancid fats are unplaitable and appear to be slightly toxic for some individuals and destructive to other factors in the food such as carotene and vitamin As and vitamin E. The preservation of a mixed food is frequently a matter largely of the prevention of fat deterioration

Prevention of Rancidity by Antioxidants. Compounds which retard the rate of development of rancidity in fats and oils are known as antioxidants. It is now generally agreed that oxidative rancidity proceeds by virtue of a chain reaction. The unsaturated fat first combines with oxygen to form a peroxide. This intermediate either spontaneously breaks down to an aldelyde or forms such a compound after interaction with water. Tho aldelyde in turn autoxidizes to a peracid, which oxidizes new, unsaturated linkages simultaneously with its transformation to an ordinary acid.

Authoridants are able to disrupt this chain reaction. Thus after the oxidant (1) is changed to a peroxide (102), the antioxidant (B) behaves as follows.

$$10_2 + B \rightarrow 10 + B0$$

 $10_2 + B \rightarrow 10 + B0$

Emulsions. When oil or liquid fat is shaken with water, it becomes finely divided and is dispersed in the water to form what is known as an emulsion. Emulsification with water alone is of course inefficient and transitory. In the presence of emulsifying agents dispersion of fat globules is more complete and hence more permanent. Among the important cmulsifving agents are soaps, proteins, bile salts, mono- and diglycerides, and such technologically useful substances as the fatty acid esters of nolvmeric glycols or sugar alcohols like Myrj, Span, Tween, etc. Emulsifying agents act by lowering the surface tension of the aqueous phase and are presumably adsorbed on the surface of the tiny oil globules to form a film which minimizes the tendency of the globules to coalcscc. A very fine powder which is net by oil but not by water will likewise form a film around oil droplets and stabilize the emulsion. Oil-in-water emulsions are also stabilized by the use of gums which increase the viscosity of the aqueous or external phase. The formation of emulsions is of importance in pharmacy, in industrial practice, and in the processes of fat digestion in the intestinal tract.

Hydrolysis of Fats. Fats may be hydrolyzed by various agents, with the liberation of fatty acids and glycerol. Such agents are superheated steam, long-continued action of air and light, bacteria, and the enzyme lipase. When boiled with alkali the fats give glycerol and the metallic salt of the fatty acid.

$\begin{array}{c} C_3H_3(\mathrm{OOCC_{15}H_{31}})_3 + 3\mathrm{NaOH} \rightarrow C_4H_3(\mathrm{OH})_3 + 3\mathrm{C_{15}H_{31}COONa} \\ \mathrm{Tripalmitin} & \mathrm{Glycerol} & \mathrm{Sodium\ paimitate} \end{array}$

This process is known as saponification, and the metallic salts of the higher fatty acids are known as soaps. The ordinary hard soaps of commerce are chiefly sodium soaps. Potassium forms soft soaps, of which green soap is an example. Calcium and magnesium form insoluble soaps. When ordinary soap is added to hard water a certam amount is used up in the precipitation of calcium and magnesium salts before the soap becomes effective. Lead salts also form insoluble soaps.

The detergent (cleansing) action of soaps is due largely to their ability to lower surface tension and thus facilitate emulsification of oily or greasy material, which can then be washed away. The commercial production of synthetic detergents has become of importance; these may be exemplified by one type which is a compound of sulfuric acid and cetyl alcohol, Chiliofil, the alcohol analogue of palmitic acid. Synthetic detergents act similarly to soaps by lowering surface tension; they are not inactivated by calcium and magnesium, however, and are therefore equally effective in hard and soft water, as the ordinary soaps are not.

Methods Used in Study of Fats. As the separation and identification of individual fats is a difficult matter, the properties commonly studied are of a general character. The saponification value of a fat is the number of milligrams of KOH required to neutralize the free or combined fatty acid in 1 g, of fat. It is determined by saponification and titration of excess alkali and is a measure of the mean molecular weight of the fatty acids present in the fat. The Reichert-Meissl number of a fat is a measure of the amount of volatile fatty acids and is determined by titration of the steam

distillate The volatile fatty acids which also include the soluble ones, are those containing ten or fewer carbon atoms. Butter is rich in these, but most butter substitutes are not. The determination of these acids is of use in detecting adulteration of butter.

The unsaturated fatty acids, of which oleic acid is an example, differ from the saturated acids in their power to take up rodinic at the double bond —CH—CII—+ I -> —CH—CHI—The rodine rature is thus a measure of the amount of unsaturated fatty acid present in a fat. The more highly unsaturated fatty acids (di, tn., and tetracthenoid) are determined by their spectrophicometric absorption characteristics.

Waxes The waxes are esters of fatty acids with monatomic alcohols Examples are spermaccti, containing chicfly the palmitate of cetyl alcohol (C₁(H₁OH)) and beesway, consisting mainly of the palmitate of myricyl alcohol (C₂(H₄OH)). They are saponified with greater difficulty than fats and are not attacked by linease.

Compound Lipides and Sterols. See Chapter 11, Nervous Tissue and Chapter 18, Bile and Liver Function.

Blological Importance of Fats. Fats serve as a storage food in plant and animal organisms and are of great importance for cellular processes, evidently along with other lipides. They are constituents of cell membranes, and are thus concerned with the phenomena of cell permeability and cell organization. The highly unsaturated fatty acids inholeic and linoleine, appear to be necessary in the due of certain animals to prevent a skin syndrome (See Chapter 33 for a further discussion of fats and fat metabolism.)

- (h) Repeat the above test using (1) 3 to 4 drops of oiele acid and (2) a little soild carhohydrate instead of olive oil. Heat vigorously and note the odor of SO; in both tubes. Organic compounds reduce KHSO, to SO;, which is often mistaken for acrolein. Even fats, if heated too vigorously, yield SO; as well as, or after, the acrolein.
- 5. Emulsification. (a) Shake up a drop of neutrain olive oil with a little water in a test tube. The fat becomes finely divided, forming an emulsion. This is not a permanent emuision since the fat separates and rises to the top upon standing.
- (h) To 5 ml, of water in a test tube add 2 or 3 drops of 0.5 per cent Na₂CO₃. Introduce into this faintly alkaline solution a drop of neutral olive oil and shake. The emulsion, while not permanent, is not so transitory as in the case of water free from sodium carbonate.
- of water free from sodium carbonate.

 (c) Repeat (b), using rancid olive oil. What sort of emulsion results? In this case the alkall combines with the free fatty acid to form soap, and this soap, being an emulsifying agent, emulsifies the fat.
- (d) Shake a drop of neutral olive oil with dilute albumin solution. What is the nature of this emuision? Examine it under the microscope.
- (e) Repect (d), using bile sait solution instead of albumin solution. Compare the stability of the emulsion with that obtained in (a). What is one of the physiological functions of the bile?
- 6. Fat Crystals. Dissolve about 40 drops of meited lard in 10 ml, of ether in a test tube, stopper ioosely with some filter paper, and allow the mixture to evaporate spontaneously until crystals begin to separate out. Transfer some of the material to a slide, examine the crystals under the microscope, and compare them with those reproduced in Figs. 34, 35, and 36.
- 7. Saponification of Boyberry Tollow." Place ahout 10 g., of hayherry tallow in a 600-ml. beaker; add about 150 ml. of distilled water and 50 ml. of 10 per cent potassium hydroxide solution (not sodium hydroxide). Boil the mixture for 10 to 15 minutes, or until saponification is complete (no oll) layer remaining on the surface). When apponification is complete, "ir remove 25 ml. of the hot soap solution, dilute it with ahout 100 ml. of distilled water, and reserve this solution for use in Exp. 8 and 9 below. To the remainder of the hot solution slowly add concentrated hydrochloric acid until the mixture is acid (about 10 ml. are required)." The free fatty acid formed will rise to the top as a clear olly layer. Now cool the solution, permitting the fatty acid to solidify and form a cake. In this instance the fatty acid is principally paimitic

¹¹ Neutral object oil may be prepared by shaking ordinary olive oil with a 10 per cent solution of sodium carbonate. This mixture should then be extracted with other and the other removed by evaporation. The residue is neutral object of

is Bayberry t dlow is derived from the fatty covering of the berries of the wax myrtle. It is therefore frequently called myrtle wax or bayberry wax.

¹³ Place 1 or 2 drops in a test tube full of hot distilled water. If saponification is complete, the soap formed will remain in solution and no oil will separate out.

[&]quot;Under some conditions a purer product is obtained if the sorp solution is cooled before precipitating the fatty acid.

acid. Remove the cake, break it into small pieces, wash it with water by decantation, and place in a beaker containing about 50 ml. of 55 per cent alcohol. Warm the mixture by placing the heaker in a vessel containing some bot water, until the paimitic acid is dissolved; then filter through a dry filter paper and allow the filtrate to cool slowly in order to obtain satisfactory crystals.

When the palmitic acid has completely crystallized, filter off the alcohol, dry the crystals between filter papers and by exposure to the air, and try the tests given in Exp. 11. Write the reactions which bave taken place in this experiment

 Salting-out of Soap. To 100 ml. of soap solution, prepared as described above, add solid sodium chloride to the point of saturation, with continual



FIG 37. PALMITIC ACID

- (c) MELTING POINT Determine the melting point of paimitic acid by any standard method. A value lower than the theoretical will be obtained because of the presence of impurities.
- (d) FORMATION OF TRANSLUCENT SPOT OF PAPER Melt a little of the fatty acid and allow a drop to fail upon a piece of ordinary writing paper. How does this compare with the action of a fat under similar circumstances?
 - (e) ACROLEIN TEST, Apply the test as given under 4, p. 106. Expiain the result.
 - (f) IODINE ABSORPTION. TEST. For directions see Exp. 14.
 - 12. Soponification of Lard. To 25 g. of lard In a flask add 75 ml. of alcoholicpotash solution and warm on a water bath until saponification is complete.
 (This point is indicated by the complete solubility of a drop of the solution
 when allowed to fall into a little distilled water.) Now transfer the solution
 from the flask to an evaporating dish containing about 100 ml. of water and
 beat on a water hath until all the alcohol has been driven off. Acidify the solution with hydrochioric acid and cool, Remove the fatty acid which rises to
 the surface, in cutralize the solution with sodium carbonate, and evaporate
 to dryness. Extract the residue with alcohol, remove the alcohol by evaporation on a water hath, and on the residue of glycerol thus obtained make the
 tests as given below.
 - 13. Glycerol. (a) TASTE What is the taste of glycerol?
 - (b) SOLUBILITY. Try the solubility of glycerol in water, alcohol, and ether.
 - (c) ACROLEIN TEST. Repeat the test as given under 4, p. 106, using 2 drops of glycerol.
 - (d) Borax Fusion Test. Fuse a paste of glycerol and powdered borax on a platinum wire and note the characteristic green flame. This color is due to the glycerol ester of boric acid.
 - (e) BENEDICT'S TEST. Add a few drops of glycerol to 5 ml, of Benedict's reagent and boil for 2 to 3 minutes. How does the result compare with the results on the sudars?
 - (i) Solution or Cu(OH), Form a little cupric hydroxide by mixing copper sulfate and sodium hydroxide. Add a little glycerol to this suspended precipitate and note what occurs. Explain.
 - 14. Iodine Absorption Test. Dissolve 5 to 10 drops of an unsaturated organic acid, e.g., oleic acid, in about 5 mi, of chloroform. Add some lifbi's lodine solution, a drop or two at a time, and shake between additions. The solution will be decolorized if unsaturated acids are present. This is due to the absorption of the lodine. The test should be controlled by shaking chloroform and lodine solution to which no acid has been added.

solution of phenosphthalein and titrate with 05 N IICl Subtract the titration value of the control from that of the unknown. One ml of the IICl is equivalent to 002805 g of kOII Calculate the number of mg of KOII required to saponify 1g of the oil

16 Determination of Iodine Absorption Number With a clean dry pipet measure 0 3 ml of oil into a dry 100 ml Erlenmeyer flask Calculate the weight of the oil from the specific gravity (For accurate work the oil must be weighed) \dd 10 ml of carhon letrachforlde and after solution of the oil add exactly 25 ml of Wijs' lodine solution " Mix well, stopper, and put in a dark place for 1 to 2 hours Transfer quantitatively to a 500 ml flask, washing out the small flask with 10 ml of 10 per cent kI solution and with water to make a volume of about 250 ml Titrate with thiosulfate to a light brown color, add starch paste and titrate to disappearance of the blue color After the blue color has disappeared from the aqueous phase, the carbon tetrachloride layer in the bottom of the flask usually contains untitrated lodine as evi denced by a pink or violet color This lodine may be brought into the aqueous layer by continuous shaking The end point of the titration is reached when both the aqueous and nonaqueous phases are completely colorless Calculate the number of centigrams of lodine absorbed by 1 g of fat This is the sodine absorption number, or the sodine value, of the oil

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See Appendix

4

Proteins: Their Composition and Hydrolysis; Amino Acids

PROTEINS

Definition. Proteins have been defined as extremely complex nitrogencontaining organic compounds which are found in all animal and vegetable cells, where they constitute a major part of the living protoplasm of those cells The Dutch physiological chemist G J Mulder, who at the suggestion of the famous chemist, Berzehus, derived the word protein from the Greek πρώτος, "first," said in 1840 "In both plants and animals a substance is contained which is produced within the former, and imparted through the food to the latter It is one of the most complicated substances, is very changeable in composition It is unquestionably the most important of all known substances in the organic kingdom Without it no life appears possible on our planet. Through its means the chief phenomena of life are produced "Protems, carbohydrates, and fats form the three great classes of foodstuffs, but the function of protein in the diet is not primarily to supply energy, as is true of the other two, but to furnish certain essential components of the living tissue of the organism itself Although plants, including many bacteria, are capable of synthesize ing proteins from simple organic and inorganic nitrogenous compounds, this ability has been lost to such an extent by the higher animals that they must depend upon preformed proteins or rather certain of their specific degradation products, the a ammo acids, for the continuance of life

The capacity of the hying organism for storing proteins is hinited and relatively small when compared to its capacity for storing carbohydrates and fats. Proteins are, however, stored under special conditions as in eggs and seeds for use by the developing or immature organism until it can obtain food from its environment. Although the carbohydrates and lipides are both essential constituents of the colloidal complex which we call protoplasm, the proteins are of paramount importance not only because of their peculiar chemical and physicochemical properties but also because they upper it occufer upon various types of cells their biological specificity. In the main, identical hipides and curbohydrates may be found in cells of both plants and anim its of widely different species, the proteins, however, are usually highly durate traite of the specific organ in which they are found

Composition the proteins differ from earboby drates and fats not only in their function in the organism, but also in elementary composition

In addition to carbon, hydrogen, and oxygen, the proteins invariably contain introgen and generally also sulfur. The percentage composition of the large number of proteins from different sources which have been studied aliab between the following rather narrow limits, C=50 to 55 per cent, C=10 to 10 to 10 per cent, C=10 to 10 to 10 per cent. Proteins have also been described which contain phosphorus, tron, copper, todine, manganese, zinc, and other elements. None of these latter elements, with the exception of iodine, has thus farbeen found as a constituent of the α amino acids, the fundamental units from which proteins are built up by the organism, and until we have more exact information concerning the composition and structure of the protein molecule, we may assume that these elements are combined with protein in some unknown way, or, as has been shown in the case of iron, that they are constituents of nonprotein substances which are combined with protein in a way which confers new and characteristic properties upon the complex formed.

stituent units, the a-amino acids; (2) partial hydrolysis to yield relatively small degradation products (di- and tripeptides, etc.) which are capable of separation and complete characterization; (3) stepwise degradation. that is, removing one amino acid after another from one end of the polypeptide chain; and (4) the study of the physicochemical. colloid-chemical. and biological reactions of the completely intact molecule as well as of the slightly changed (denatured) molecule. Until comparatively recently the method of complete hydrolysis was the one most commonly employed. This procedure is of great importance since it yields the individual units from which the protein is formed. However, just as elementary analysis of the less complex organic molecules tells us the percentage composition of the substance under examination without indicating the atomic arrangement within the molecule, so complete hydrolysis of a protein reyeals the quantities of the various constituent amino acids without giving much indication as to their special arrangement within the giant moleculc. In order to reveal the great individual differences among proteins and to gain an insight into their specific biological functions, such as the hormonal action of insulin, the enzymatic effects of pepsiu, the role of myosiu in muscle contractiou, the pathogenicity of tobacco mosaic virus, etc., all four modes of study must be used. The work of the chemist, hiologist, physicist, and analyst must be known and appreciated by each member of the team hefore the problem of the complete structure of even one protein is to he solved.

Hydrolysis. Hydrolysis of proteins may be effected: (1) By hoiling with mineral acids or strong alkahes, at atmospheric or increased pressures, (2) hy treatment with certain long-chain sulfonic acids (cetylsulfonic acid, diphenylbenzenesulfonic acid, etc.), and (3) hy digestion with proteolytic enzymes. Commonly, hydrolysis of a protein is carried out hy boiling it with five to ten times its weight of 6 N hydrochloric acid or 8 N sulfuric acid for 6 to 24 hours. Under special circumstances other reagents, such as hydriodic acid, oxalic acid, 5 N sodium hydroxide, hot saturated harium hydroxide, or a mixture of formic and hydrochloric acids, can he used. Acid hydrolysis, especially if the protein contains carholydrate, usually results in the complete destruction of the amino acid tryptophan, and may result in the partial decomposition of certain other amino acids. The prolonged heating necessary to effect complete hydrolysis hy strong alkalies does not affect tryptophan, but results in the partial or complete destruction of cysteine, cystine, and arginine, and in the racemization and consequent loss of optical activity of all the amino acids. Enzymatic hydrolysis has none of these disadvantages of acid or alkaline hydrolysis but it is very time-consuming and is seldom complete. Hydrolysis of proteins with the above-mentioned sulfonic acids requires further investigation.

Hydrolysis of the protein molecule by any of the methods above leads to the formation of a series of ill-defined fragments of decreasing complexity known as proteoses, peptones, and polypeptides, the final products being ammo acids. Certain ammo acids are split off early in the hydrolytic process. This riberation of ammo acids continues until the larger intermediate fragments have all been reduced to these simpler complete the process.

pounds Physically the hydrolysis of proteins consists in a breaking down of the large, colloidal, nondifficultie complexes into a series of fragments which the colloidal character becomes less and less pronounced, untilinally only the simple, crystalloidal and diffusible amino acids remain. Since the amino acids represent, therefore, the "building stones" of the protein molecule we may well begin the study of that molecule by a consideration of the structure and reactions of these fundamental units

AMINO ACIDS

The amno acids thus far isolated from protein hydroly zates are a mino acids—that is they have an amino (NHs) group attached to the same carbon atom that holds the carboxyl (COOH) group. Their general for mula' is therefore

Individual amino acids differ in the character of the radical R attached to the a carbon atom

CLASSIFICATION

The amino acids may be conveniently classified according to the number of their amino and carboxyl groups, as follows (1) Neutral amino acids containing one amino and one carboxyl group, (2) acide amino acids, containing an excess of carboxyl groups and (3) hasic amino acid-containing an excess of basic introjen. The amino acids in each group may be further subdivided according to whether the radical R in the general formula represents an alphatic aromatic or heterocyclic nucleus.

NEUTRAL MINO ACIDS

These acids are often referred to as monoamino-monocarboxylic acids because they contain one amino and one carboxyl group. Solutions of these acids react essentially neutral. These acids form the largest group in the protein molecule and most of them may be separated in one fraction from the products of protein hydrolysis. Because of their great chemical and physical similarities, the isolation of the individual acids in this fraction presents experimental difficulties.

2. Alanine, C₃H₇O₂N (α-aminopropionic acid)

3. Serine, C₃H₇O₅N(β-hydroxy-α-aminopropionic acid or β-hydroxy-alanine)

4. Threonine, C4H9O2N (α-amino-β-hydroxy-n-butyric acid)

5. Valine, $C_5H_{11}O_2N$ (α -aminoisovaleric acid or β,β -dimethylalanine)

6. Leucine, $C_6H_{13}O_2N$ (α -aminoisocaproic acid or β -isopropylalanine)

7. Isoleucine, $C_6H_{13}O_2N$ (β -methyl- α -aminovalene acid or β -methyl- β -ethylalanine)

B. AROMATIC AMINO ACIDS

8. Phenylalanine, $\rm C_9H_{11}O_2N$ $(\beta\text{-phenyl-}\alpha\text{-aminopropionic acid or }\beta\text{-phenylalanine})$

 Tyrosine, C₂H₁₁O₂N (β-parally droxyphenyl-α-ammopropionic acid or β-parally droxyphenylalanine)

C SULFUR-CONTAINING AMINO ACIDS

Cysteine, C₁H₇O₂NS (β-thiol-α-aminopropionic acid)

Cystine, C₄H₁₂O₄N₂S₂ (di-(β-thiol-α-aminopropionic acid))

12. Methionine, C4H11O2NS (7-methylthiol-a-amino-n-butyric acid)

D HETEROCYCLIC AMINO ACIDS

13. Tryptophan, $C_{11}\text{II}_{12}\text{O}_2\text{N}_2$ (\$\beta\$-3-indole-\alpha\$-aminopropionic acid of \$\beta\$-indolealamne)

² Cystine is included here among the monoamino-monocarboxylic acids because it contains equal numbers of amine and earboxyl groups

14 Proline, C5H9O2N (pyrrolidine-2 carbovylic acid)

15. Hydroxyproline, $C_2H_9O_3N$ (oxyproline or 4 hydroxypyriolidine 2 carboxylic acid)

ACIDIC AMINO ACIDS

These acids often referred to as monoamine-dicarboxylic acids contain more carboxyl than uning groups and are therefore acid in reaction. The acidic nature of glutamic acid and aspartic acid allows their ready separation from the other components of protein hydrolyzates.

16 Aspartic acid, C.H.O.N (α aminosuccinic acid)

17 Glutamic acid, C.H.O.N (α ammoglut iric acid)

BASIC MINO ACIDS

These acids are predominantly basic in reaction and are precipitated from protein hydrolyzates by the addition of phosphotungsite acid or they may be selectively adsorbed on neak cation exchange resins

I Proline and cystine are also aree p tated to a certain extent by pl osphotungstic acid.

 IlistidIne, C₆II₂O₂N₂ (β-imidazole-α-aminopropionic acid or imidazolealanine)

Arginine, C₄Π₁₄O₂N₄ (δ-guanidino-α-aminovaleric acid)

20. Lyslne, C. H 14O2N2 (α-e-diaminocaproic acid)

Hydroxylysine, C₄H₁₅O₂N₂ (α-ε-diamino-δ-hydroxycaproic acid)-

22. Citrulline, C₆H₁₂O₂N₂ (δ-carbamino-α-aminovaleric acid)

Other Hydroiysis Products of Proteins. Most proteins contain practically all of the amino aculs listed above Occasionally we find that certain amino acids are either entirely absent in a particular protein, or else present in amounts too small to be detected by the available methods of analysis. Thus the protein zein, from maize, contains no lysine or glycine, gelatin yields no tryptophan, and insulin contains no methionine. In other cases we may find that certain specialized proteins contain amino acids not found in any other proteins. The most noteworthy example of this is a globulin obtained from the thyroid gland, thyroglobulin, which normally contains the iodinated tyrosine derivatives, monoiodotyrosine;

3, 5-dnodotyrosme, or nodogorgone and, and β -1'-hydroxyphenyl-3',5',3, 5-tetranodotyrosme, or thyroxme. Interestingly enough the hromine analogues of nodotyrosmes—namely monobromotyrosme and dibromotyrosme—have been found in a number of Mediterranean corals and Gorgoma.

Other Porifers and coclenterates yield monoiodotyrosine, duodotyrosine, and even traces of thyroxine

Occasionally substances known as diketopiperazines, which are anhydrides formed by the condensation of two molecules of amino acid, have been found in protein hydrolyzates. It is probable that these are formed during the treatment of the hydrolyzate

In addition to the above-mentioned amino acids, there are other less well-recognized amino acids in many peptide molecules. Thus a-c-diaminopimehe acid his been found in the insoluble proteins of the microorganism Corynebaclerium diphtheriae, and iodinated histidine has been reported to be present in pancreatic digests of thyroglobulin.

Acid hydrolysis of proteins also liberates variable quantities of ammonia which come primarily from the hydrolysis of the acid anide groups of asparagine and glutamine, for it is now known that a portion of these dicarbovylic runno acids exists in the peptide chain with the "omega" carbovyl group aminated A smaller portion of the ammonia presumably comes from the hydrolytic decomposition of certain amino acids such as cysteme, serine, threonine, etc. If ilk-thine hydrolysis is employed, ammonia is formed from glutamine, asparagine, cystine, cysteme, serine, threonine, and argimine. The last-named substance breaks down to yield ornithine and two molecules of animonia.

Hydrolysis of proteins also yields small quantities of carbon dioxide, hydrogen sulfide, and free sulfur. These products arise from the destructive decomposition of the more Lable ammo acids, especially cysteine Many proteins (serum albumin, egg albumin, thrombin) yield varying amounts of carbohydrate. These carbohydrates appear to be polysactianties composed of glucosimine and hevose (miniose, galactose). It is not known whether the cirbohydrates are an integral part of the protein molecule or whether the protein is composed of a carbohydrate-free portion admixed with a small quantity of a carbohydrate-rich protein (glycoprotein) such as seromucoid, seroglycoid, ovonucoid, etc.

Determination of Amino Acids. The quantitative determination of each of the component amino acids produced during by drolysis of a protein is a problem of greatest importance since any theory of protein structure must rest ultimately on an exact knowledge of all the units contained in the molecule. This problem is still far from solved since it presents man serious difficulties, not least of which is the preliminary by drolysis of the protein molecule into its component amino acids without causing decomposition during the bytorysis.

APPROXIMATE PERCENTAGE COMPOSITION OF SELECTED PLANT PROTEINS*

	Gliadin	Zein	Corn Gluten*	II heat Gluten*	Polato Meal*	Peanul Meal*	Sojbean Meal*
∖itrogen	17 66	16 2	12 7			10.8	7 9
Sulfur	1 24	0 ა2	15			10 5	
Arginine	27	17	3 1	3 9	50	10 6	7 3
Histoline	23 1	1 3	2 1	2 2	2 2	2 1	29
1 ysine	īí	0 0	1 5	19	83	3 0	6.8
Tyrosine	3 2	53	63	38	2.5	4.4	40
Try ptophan	0.6	0.1	0.6	0.8	2 1	10	14
Phenylalanine	6.9	6 2	6 6	5 5	54	5 1	5 3
Cystine	26	0.8	1.5	2 4	1 3	10	19
Methionine	17	24	2 5	17	20	10	1 7
Threonine	2 1	3 5	4.0	2 5	6.9	16	3 9
Senne	4 9	83		4 0	26	66	4 2
Laucine	65	23 7	16 0	7.0	9 6	6.7	8.0
Isoleucine	5 4	73	5 1	1 4 2			60
Valine	2 7	3 5	37	4 1	3 7 5 3	4 0	5 3
Glutamic acid	45 7	26 9	21,	35 0		·	18 4
Aspartic acid	1 3	66		3 8	7 4 11 5	17 4 15 1	3 7
Glycine	<0 2	04					1
Alanine	21	116	4 3	3 3	19	1 5	40
	2 1	1116	i	2 7	6 1	4 2	3 3
Proline	13 4	10 5	i	14 1	3 0	5 2	5

Calculated in g of amino acid per 160 g. of ritrogen

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B Van Slyke's Nitrogen-Distribution Method When an aliphatic primary amine is treated with nitrous acid, its amino nitrogen is quantitatively converted into free nitrogen

$R NH_2 + HONO \rightarrow R OH + H_2O + N_2 \uparrow$

By the use of this reaction, supplemented by other procedures as indicated in the accompanying diagram, Van Slyke developed a method for the determination of the distribution of amino and nonamino introgen in the protein together with fairly accurate determinations of cystine, argmine, lysine, and histidine

DISTRIBUTION OF PROTEIN NITROGEN BY VAN SLYKE METHOD

- I Amide Nitrogen (Distilled off as ammonia after neutralization of hydrolysis mixture with magnesium oxide) a Cystine (Contains only amino N Esti mated from sulfur determinations) II Basic Nitrogen b Lysine (Contains only amino N)
 (Precipitated) a Arginme (Three quarters of nitrogen is nonamino N Half of nitrogen liberated as by phospho-) Total Natrogen ammonia by boiling with strong NaOH) tungstic acid)/ (Kieldahl ded Histidine (Two thirds of nitrogen is non termination (/ omma on portion of Monoamino N (Neutral and hydrolysis acid amino acid N One-half mixture) III Monoamino Nitrogen + Non of tryptophan N Deter mined by Van Slyke amino Nitrogen (Kieldahl determination on filtrate from method) precipitated phosphotung b Nonamino N (Proline, hy droxyproline and one half states) tryptophan N Difference between III and IIIa)
- C Determination of Individual Amino Acids Differences in the chemical nature of the various amino acids have been utilized to permit the estimation by chemical means of the quantities of individual amino acids present in a protein hydrolyzate. The principal methods in use at the present time are outlined below
- I Glycine in a protein hydrolyzate is oxidized with ninhydrin (triketohydrindene hydrate) to yield formaldehyde
 - 2 Alanine is oxidized with ninhydrin to yield acetaldehyde
- 3 Serine is oxidized with periodic acid to yield formaldehyde which is readily determined
 - 4 Threonine is oxidized with periodic acid to yield acetaldely do
- 5 Phenylalanme is first nitrated to yield 3,4-dimtrophenylalanme, this is reduced to the nitroso compound, which has a purple color in alkaline solution
- 6 Tyrosme, like other 3,5 unsubstituted phenols, gives an intense red color when treated with mercury salts and nitrous acid (Millon-Nasse reaction)
 - 7 1ry ptophan gives various red and purple compounds when con-

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. Calculated in g of amino acid per 16 0 g of nitrugen

densed with aromatic or aliphatic midely less in the presence of concentrated hydrochloric or sulfuric icil (Hopkins Cole, Voiscuet Rhode reactions)

8 Cystine is reduced to cysteme. The latter is capable of reducing phospho-18-tingstic real to viel the deep-blue lower oxides of tingstin 9 Methomnic is treated with sodium introprusside to give a colored

complex 10 Aspartic read is first procipitated as the calcium with and then is isolated as copper aspart ite

11 Glutanue acid is oxidized with chlor nume-1 to β-ryanopropiorie reid. The latter is hydrolyzed to succume acid which is then determined

12 Histidine is precipitated as the silver salt at pH 74 and the isolated as histidine nitramlate Nitramlie und is 2,5-dihydroxy-36-

dinitro-p-benzogimone 13 Argumus is first precipitated by silver intrate and bary to at strongly alkalme reaction and then isnisted as pure arginine 2 I-dimitro-1 naph

thol 7 sulfonate (arginue miniolinamente)

14 I same can be determined in the intact protein by virtue of the fact that almost all of the free ammo groups in a protein are tho e immo groups of lysme Lysme is isolated from a protein hydrolyzate previously freed of histidine and argmine by precipitation with phospho-21 tunistic acid and subsciment isolation as lysine merate

10 Hydroxylysine is precipitated with phosphotungstie acid. On treat ment with periodic acid in alkalino solution one molecule of hydroxylv me

yields one molecule of formaldeliade and one of ammonia

D The Isotope Dilution Method of Usern and of Schoenheimer and Rittenberg 1 compound which has an abnormal isotope content is in separable from its normal analogue by the usual laboratory procedures Thus if an amino acid containing a known excess of an isotope is added to a protein hydrolyzate the quantity of the amino acid present in the hydrolyzate can be calcolated by a determination of the isotope content of the isolated amino acid. The yield of amino acid actually isolated is of no importance in this method

Isotopic-Carrier Method 1 mixture of amino acids is treated with a reagent containing stable or radioactive isotopes to form quantitatively a stable derivative of the desired constituent. Then an overwhelming ev cess W of the unlabeled derivative (the earrier) is added. The carrier is

isolated and purified to constant isotope concentration C.

If C is the isotopic concentration of the pore isotopic derivative prepared with the same reagent the quantity of derivative present 18 W X C /C Thus one isotopic reagent soffices for the analysis of man)

compounds

E Microbiological Procedures Various microorganisms such as Lactobacillus arabinosus etc require certain amino acids for normal growth (see p 1062) Preparation of synthetic media lacking in only one of these amino acids permits the determination of the specific amino acid in the unknown solution Growth of the microorganism on the synthetic medium supplemented by the solution undergoing analysis is compared with that obtained in the presence of known amounts of added amino acid In

other cases, mutants of the mold Neurospora crassa have been developed which require only one amino acid for growth. A number of enzymes have heen separated from plant and animal sources, each of which decomposes only a single amino acid in a mixture. Thus arginine may be determined by the action of arginase, which liberates ammonia; tyrosine, histidine, lysine, and glutamic acid are each decarboxylated by specific enzymes. Quautitative determination of the carbon dioxide thus liberated is a measure of the amino acid present.

F. Chromatographic and Ion-Exchange Methods. The powerful analytical tools of paper chromatography and ion-exchange chromatography have not only been applied to the qualitative and quantitative determination of all the amino acids which commonly occur in protein hydrolyzates but have enabled investigators to find hitherto unsuspected amino acids, e.g., monoiodotyrosine, \$\varepsilon\$-hydroryphenyl-trinodotyrosine, diaminopimelic acid, etc. Plate I shows the separation of the amino acids in a protein (keratin from wool fiber) hydrolyzate on a two-dimensional paper chromatogram after staining with ninhydrin. This type of chromatogram is excellent for the qualitative identification of amino acids and it may be employed, under proper conditions, for their quantitative estimation. The quantitative or, more usually, semiquantitative estimation is based on the observation that the maximum color density of each spot is proportional to the concentration of material at that spot

The amino acids may also be separated one from another by first adsorbing the mixture on a column of a polystyrene cation-exchange resin. The iudividual amino acids are then eluted from the resin with huffers of progressively increasing pH. Many small fractions of the cluate are collected separately and each fraction is analyzed for its amino acid content by heating with ninhydrin. From the data thus obtained, the quantity aud identity of each amino acid in the mixture may be computed.

G. Electrolytic Separations. The α -amino acids may be divided into three groups—neutral, acidic, basic—by virtue of their charge in solutions of definite hydrogen-ion concentrations. Electrolytic separations are

employed preliminary to paper chromatography.

General Properties of Amino Acids. The amino acids derived from proteins are all a-amino acids. The a carbon atom in all the acids, with the exception of glycine, is asymmetric, so that these acids are all optically active. They are white crystalline substances, the crystal form heing characteristic for each acid. They are all soluble in water, except cystine and tyrosine, the latter being more soluble in hot thau in cold water. With the exception of proline, they are all insoluble in alcohol, and all are insoluble in ether. They are, as a rule, all soluble in solutions of strong acids and bases. They are not precipitated by ammonium sulfate or sodium chloride, but with the exception of proline are precipitated by alcohol. They form crystalline salts with metallic bases and with mineral acids. Many of the amino acids (such as glycine, alanine, serine, and proline) have a sweet taste; some, like tryptophan and leucine, are tasteless; others, like arginine, are bitter. The amino acids are all amphoteric, ionizing both as acids and as bases by virtue of their carbovyl and amino groups, thus forming salts with labiles and with acids.

Spatial Configuration and Optical Activity of Amino Acids. The amino acids have the following general structure

In this structure the α carbon atom is asymmetrie—i e, there are for different substituent groups. I has all the amino acids except given (where R = H) are optically active, and are capable of existing in two different spatial forms, which are mirror images of one another, will equal and opposite optical rotatory power, and which differ solely in the arrangement of the substituent groups in space around the α carbon atom. These two different spatial arrangements are known respectively as the p and L configurations

The configuration of a particular amino acid is based on its spatial relationship to an arbitrary reference compound, just as the configuration of p-glucose is hased on that of dextrorotatory p-glyceraldehyde (& Chapter 2) This relationship, for the amino acids, is illustrated as follows:

Thus the naturally occurring amino acid alanine, which is dextrorola tory, may be suitable means be related structurally to either dextrorola tory lactic acid or les orotatory glyceraldehyde. Both of these have the same configuration, which in this case is known as the Loonfiguration when referring to amino acids As with the sugars there is no necessary agreement between configuration and the direction of optical rotation, to indicate the latter, the signs (+) for dextrorotatory and (-) for levorotatory are used. Naturally occurring alanine is therefore L(+)-alanine, and the direction of the configuration and the direction of the configuration and the direction of optical rotation, to rotatory are used. Naturally occurring alanine is therefore L(+)-alanine, and the direction of the configuration of the configuration of the configuration of the configuration.

The spatial configuration of an amino acid appears to have important physiological significance. For instance, certain (but not all) of the amino acids cannot be utilized by the animal body if the configuration is opposite to that found naturally. It is an interesting fact that the bulk of the amino acids found in nature either free or as part of proteins, have the L configuration Certain a amino acids have however been isolated from natural sources thus a glutamic acid is a major constituent of the capsule of Bacillus anthraces and related microrganisms the naturally occurring acids and o value is a decomposition product of the penicillus. The significance of these stereochemical findings remains to be elucidated.

With alkalies, such as sodium hydroxide, the ionization proceeds to the right and leads to the formation of a sodium salt of the amino acid, which is ionized. This gives a positive sodium ion and a negative amino acid ion

In acid solutions, therefore, the amino acid earnes a positive charge and in an electrical field migrates to the cathode, but in alkaline solutions it carries a negative charge and migrates to the anode For every amino acid there is a definite hydrogen-ion concentration, specific for that acid, at which the degrees of acid and basic ionization are equal. At this particular pHI, known as the isoelectric point, the amino acid is electrically neutral and will not migrate in an electrical field.

In past years there has been a great accumulation of experimental evidence indicating that this electrically neutral form of the amino acid consists of a mixture of undissociated molecules and a tautomeric form, known as a zwitterion, in which both the amino and earboxyl groups

are ionized to the same extent. In solutions of aliphatic amino acids, more than 99 per cent of the molecules are present in the form of zwitterions, in the aromatic acids the two forms are present in approximately equal amounts. The addition of acid to the zwitterion suppresses the ionization.

of the carboxyl group, thus leaving a positively charged ion which is free to form salts with the acid Sumlarly alkali suppresses the ionization of the amino group, thus taving a negatively charged ion which can form salts with the base. Thus the effects of adding acid or alkali to the amino

acid are the same regardless of whether the neutral amino acid is considered to be an undissociated molecule or a zwitterion

These reactions of amino acids with acids and with bases are of great importance in protein chemistry since, as we shall see in Chapter 5 a great many of the physicochemical reactions of proteins are explainable on the basis that the protein molecule contains a definite number of free amino and free carboxyl groups the exact number of each being characteristic of each particular protein. Depending upon the pH of the solution therefore, proteins combine with acids and bases and carry a preponder ance of either positive or negative charges, or behave as though they were electrically neutral

REACTIONS OF AMINO ACIDS WITH NITROUS ACID The amino acids as their general formula indicates are primary amines and like all such amines yield nitrogen when treated with nitrous acid. This reaction

$$\begin{array}{c} H \\ R-C-COOH + HNO \rightarrow R-C-COOH + N_2 + H_2O \\ NH_2 \end{array}$$

forms the basis for Van Slyko's method for the determination of free amino groups, as has already been indicated. For this purpose use is made of a specially devised apparatus in which the nitrogen gas evolved during the reaction is collected and its volume measured. This reaction is important for the determination of free amino groups in amino acids or mixtures of amino acids, and is also used in estimating the amounts of amino acids in biological fluids, such as blood. It is also used to determine the percentage of the total introgen of the protein that is present in the form of free amino introgen. Since the latter increases during acid or enzymatic hydrolysis the method is also of great value in determining the rate and extent of protein degradation by any of the hydrolytic agents.

REACTIONS OF AMINO ACIDS WITH NINHYDNIN The amino acids in general react with the compound mulhydrin (C₄H₄ (CO)₂ H₂O, triketohy drindene hydrate) to yield earlion dioxide aminoma and usually (but not always) an aldehyde containing one fewer carbon atom than the original mino acid

$$\begin{array}{c} NII_{2} \\ R - C - COOH \rightarrow R - C = 0 + NII_{1} + CO_{2} \\ II & H \end{array}$$

I his reaction has been made the basis for several different types of quantitative methods for the determination of amino acids. These are based upon (1) the color change which results from the reaction (see pp. 18 and 172), (2) determination of the aminoma produced, and (3) measurement by gasometric means of the curbon dioxide evolved. This latter procedure has been accurately established by Van Slyke and his associates as possibly the most satisfactory method available for the determination of a unino introgen (in terms of CO-), exceeding the nutrous acid method

in specificity for this purpo e. For a description of this method as applied to the determination of amino acid nitrogen in urine, see p. 892

REACTIONS OF AMINO ACIDS WITH FORMALDEIN DE The carboxyl group of the simple amino acids is not readily titratable with alkali under ordinary conditions presumably because of the influence of the neighboring amino group. In 1899 Schiff observed that in the presence of formaldehyde the amino acids became as readily titratable as any simple organic acid. Sprensen formulated the reaction between amino acids and formal dehyde as follows.

$$R-CH-COOH + H-C-H \rightarrow R-CH-COOH + H_2O$$
 H_2
 O
 CH_2
 CH_2

There is considerable evidence against this view however and Harris believes that only an amino acid formaldehyde complex is formed

Others have suggested that HCHO reacts with amino acids to give monor dimethylol derivatives

The presence of formaldehyde decreases the basicity of the amino group permitting the carboxyl group to exert its maximum acidity. This acidit may then be titrated with standard sodium by drovade using phenol phthalein as indicator. This reaction forms the basis for Sprensen's for litration method for the estimation of free carhoxyl groups in amino acids and mixtures of amino acids. It is also widely used for determining yais of proteins

Reactions of Anino Acids with Amno Acids Many of the amino acids readily form anhydrides when their solutions are evaporated. This reaction in volves a condensation between the amino group of each molecule with the carboxyl group of the other the resulting compounds being known as diletopiperazines. In 1901 Fischer and Fourneau subjected

chycine anhydride to weak hydrolysis with acids and obtained a compound glycyl-glycine in which the amino group of one acid was combined

with the carboxyl group of the other This was the starting point of a great

many researches whereby Fischer and his co-workers, using various derivatives of amino acids, prepared a large number of similar compounds between amino acids, called dipeptides. Although the details of these methods would be out of place in a book of this character, the reactions involved were chosen so that the amino group of one acid always combined with the carboxyl group of the other resulting in the so-called peptide linkage. Since these dipeptides still contained a free amino and a

earhoxyl group, by the use of various migenious methods Fischer was ablo to lengthen these chains of amino acids, forming polypeptides containing as many as 18 acids. According to the modern conceptions of protein structure, the protein molecule consists in large part, at least, of amino acids linked together through their amino and carboxyl groups. The evidence for this point of view will be discussed more fully in the section on the structure of the protein molecule in Chapter 5.

DISCUSSION OF THE INDIVIDUAL AMINO ACIDS

Glycine, C2H3O2N (aminoocetic acid, glycocoll).

Glyeme was the <u>first amon and to</u> be isolated as a primary decomposition product of the proteins. Glyeme is prepared synthetically by the action of ammonia on monochlorocette acid. It crystallizes in rather large colorless monochine crystals usually in four-sided prisms. Glyeme pos-cases no asymmetric earbon atom and is therefore not optically active.

In addition to being present in many animal proteins, glycine is found in the body as a constituent of various nonprotein nitrogenous compounds it is a component of (1) glutathone, a tripeptide of glutamic acid, cysteine, and glycine (see p 116), (2) glycochobe acid, a compound

of glycine and choice and, found in the bile (see p. 410); and (3) hippund and, or benzoyl-glycine, which is found in the nrine after the ingestion of benzoic acid or compounds which give rise to benzoic acid in metabo-



Fig. 38, GLYCINE (SYNTHETIC)

See also Fig. 103

From Keenan J. Bool. Chem., 62, 163 (1924)

lism. The reaction leading to the formation of hippuric acid from benzoic acid and glycine is as follows:

 $\begin{array}{c} C_{4}\Pi_{2}COOH + \Pi_{2}N - C\Pi_{2}COOH \rightarrow C_{4}\Pi_{3}CO - NH - C\Pi_{2}COOH + \Pi_{2}O \\ \text{Benzoic acid} & \text{Hippuric acid} \end{array}$

Alanme is best prepared from silk in which it occurs to the extent of approximately 25 per cent. It has a sweetish taste, and is dextrorotatory



From Leenan J Biol Chen 62 163 (1974)

L(-)-Serine, C₃H₂O₂N (β hydroxy-α-aminopropionic acid).

Serine crystallizes from water solution as thin, irregular plates and has a sweet taste. It melts at about 245° and is soluble in 23 times its weight of water at room temperature.

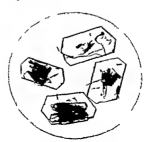


Fig 40 State.
From heenan J B of Chem 62, 163 (1974).

L(-)-Threonine, C₄H₂O₂N (α-amino-β-hydroxy-n-butyric acid).

The α carbon atom of naturally occurring threonine has the L configuration, as shown. When, however, the configuration of the molecule as a whole is considered, it is seen to resemble that of the sugar D-threose (whence the name) so that this amino acid is sometimes referred to as D(-)-threonine.

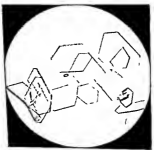


FIG 41 THREONINE

From McCor, Meyer, and Rose J Bud Chem 112, 288 (1935) L(+)-Vallne, C.H.11O2N (a-aminoisovaleric acid),

Valine occurs in casem and in egg albumin to the extent of 6 to 7 per cent. It is made synthetically by the interaction of ammonia and α -bro-monous aleric acid under pressure in which process the bromne is split off as IIBr and the amino group takes the α -position originally occupied by the bromnes.

L(-)-Leucine, C₄H₁₀O₃N (a-aminoisocaproic acid).
CH₂ NH₃

Leucine is found in almost all proteins. Free leucine has been found pathologically in urine (in acute yellow atrophy of the liver, phosphorus poisoning, and acute febrile conditions). Pure leucine crystallizes in shiny, white, extremely thin plates. Inactive leucine tastes slightly sweet, $\mathfrak{v}(+)$ -leucine tastes quite sweet, and $\mathfrak{v}(-)$ -leucine slightly bitter.



FIG 42 LEUCINE.

For the material from which the above crystals were prepared, as well as those reproduced in Figs. 43 to 45 and 47 to 48, the authors are indebted to the late Dr. Thomas B. Osborne

EXPERIMENTS ON LEUCINE

Preparation of Leucine. In a 2-liter flask, place I liter of defibrinated blood and gradually add 150 ml. of concentrated sulfuric acid, shaking well during the additions. Boil on a sand bath for 12 to 14 hours, being careful to shake continually until it boils evenly. To the hot liquid add a solution of barium hydroxide until the mixture is alkaline to litmus. Filter on a Buchner funnel. Make the filtrate acid to litmus with dijute H₃SO₄, decolorize with 20g. of activated charcoal, filter, and concentrate in a porcelain dish over an open flame to 500 ml. and filter again. Make the filtrate faintly alkaline to litmus through the addition of ammonia and concentrate on a boiling water bath until a crystalline mass forms on top of the liquid. Cool for 24 hours in the icebox. Filter on a Buchner funnel and press the water out of the crystals. Recrystallize from 70 per cent afcohol.

Make the following tests upon the leucine crystals already prepared, or upon some pure leucine furnished by the instructor.

I, 2 and 3. Do these experiments according to the directions given for Tyrosine (p. 138),

L(+)-Isoleucine, C₆II₁₃O₂N (β-methyl-α-aminavaleric acid).

Isoleucine has a bitter taste and it crystallizes in plates like leucine.

L(-)-Phenylalanine, C₂H₁₁O₂N (β-phenyl-α-aminopropionic)
acid).

Phenylalanine is easily soluble in hot water, quite insoluble in colwater, and only slightly soluble in ethyl and methyl alcohols. It has bitter taste



FIG 43 PHENYLALANINE

L(—)-Tyrosine, C₁H₁₁O₂N (β-parahydroxyphenyl-α-aminopto-pionic acid).

$$HO \longrightarrow CH^{2} - COOH$$

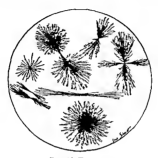
Of all the amino acids tyrosine is the least soluble in water (1 part in 2500 of water at room temperature). It is insoluble in ether, alcohol, acetone, and glacial acetic acid, but readily soluble in dilute alkalis and acids 1t crystallizes from water in sheathlike groups of fine white needles. Tyrosine is found in most proteins but is virtually absent in gelatin (0.3 reaction). It may be detected in the faintest traces by means of Millon's reaction. (See p. 169.)

EXPERIMENTS ON TYROSINE

Preparation of Tyrotine Introduce 200 g of commercial caseln into a 3-liter pyrex flask, add a liter of cold water, and shake well With continuous shakings slowly add a liter of boiling water to this mixture. Now add 40 ml. of 20 per

cent sodium hydroxide to dissolve the casein, and adjust the reaction of the casein solution to pli 8 by the addition of normal sodium hydroxide 4

Preserve the casein solution and diminish oxidase action by adding 15 ml of toluol and 2 g of sodium fluoride dissolved in about 10 ml of hot water Shake well, add trypsin in the form of some commercial preparation, and, after mixing the contents of the flash thoroughly, stopper the flash and place it in an incubator at 38° to 40° Shake the flash thoroughly every day, without removing the stopper, and at the end of the fourth day add an addi-



Tto 44 TYROSINE

tional quantity of trypsin in one of the forms mentioned After permitting the digestion to continue for a second period of four days, remove the flask from the incubator, allow it to stand at room temperature for at least 24 hours, then filter off the precipitate of tyrosine, undicested casein, etc. Treat the residue with dilute sulfuric acid (5 ml of concentrated sulfuric acid in 250 ml of water) to dissolve the tyrosine, filter through a pleated paper, add 10 ml of concentrated ammonium hydroxide to the filtrate, and heat on a boiling water bath. The solution, which should now he acid to litmus, is carefully neutralized by the addition of ammonium hydroxide and allow to cool Tyrosine, contaminated with more or less calcium phos phate, should crystallize out Filter off the tyrosine by suction, suspend it in 300 ml of water in a flash, heat to bosting, add 5 ml of concentrated ammonium hydroxide, and boil for 15 minutes Filter off the insoluble calcium phosphate, neutralize the tyrosine filtrate with 5 per cent H SO, and allow it to stand Filter off the tyrosine crystals by suction, wash well with cold water and alcohol in turn, and dry in an oven or incubator if the tyrosine crystals are not well formed (see Fig. 44) they may be recrystallized from hot water (solubility 1 154)

⁴ To do this take 10 mi of the casem mixture add t0 drops of cresol red and titrate with 22 N 'addl (using a uncrof uret or a 1m | prept graduated in 00 ml) until a reddish purple color is obtained. Multiply this ittration volume by 40 and add this quantity of nor mi sodium by droude to the casem solution in the prex flash. Shake the mixture at frequent intervals after the by droude 28 added. The reaction should now be acid to phenol phthale a and alkaline to creed red.

If pure try pain is not available the enzyme may be added in the form of pancreating. This filtration can be speeded up by the use of a diatomaceous-earth filter aid

Make the following tests with the tyrosine crystals prepared in the above experiment or upon some pure tyrosine furnished by the instructor

- I Microscopic Examination Place a minute crystal of tyrosine on a slide add a drop of water cover with a cover glass and examine microscopically Nowrun more water under the cover glass and warm in a Bunsen flame until the tyrosine has dissoived Milow the solution to cool slowly, then examinagan microscopically, and compare the crystals with those shown in Fig. 44
- 2 Solubility Try the solubility of very small amounts of tyrosine in cole and hot water cold and hot 95 per cent alcohol dilute NII OII, dliute KOII and dilute IICI
 - 3 Sublimation Place a little tyrosine in a dry test tube, heat gently an notice that the material does not sublime. How does this compare wit leucine?
 - 4 Hofmann s Reaction This is the name given to Millon s reaction wheemployed to detect tyrosine Add about 3 ml of water and a few drops (avoid an excess) of Vililon s reagent to a little tyrosine in a test tube Upon dissolving the tyrosine by heat the solution gradually darkens ond may assume a dark red color What group does this test show to be present in tyrosine?
 - 5 Sulfura Acid Test (Fira) Warm a little tyrosine on a watch glass on a boiling water bath for 20 minutes with 3 to 5 drops of concentrated II;50. Tyrosine sulfura caid is formed in the process Cool the solution and wash it into a small beaker with water Now slowly add CaCO₂ in substance with string until the reaction of the solution is no longer and Filter concentrate the filtrate and add it to a few drops (avoid an excess) of very dilute neutral ferric chloride A purple or violet color due to the formation of the ferric sail of tyrosine sulfura caid is produced. This is one of the most satisfactory tests for the identification of tyrosine.

$$2 \text{HOC}_6 \text{H}_6 \text{CH}_2 \text{CHNH}_2 \text{COOH} + 2 \text{I}_2 \rightarrow 2 \text{HO} \underbrace{\hspace{1cm} \text{I}}_{\text{1}} \text{CH}_2 \text{CHNH}_2 \text{COOH}$$

$$\text{Tyrosine} \qquad \qquad 3,5 \text{-Dilodotyrosine}$$

$$\rightarrow \text{HO} \underbrace{\hspace{1cm} \text{I}}_{\text{2}} \text{CH}_2 \text{CHNH}_2 \text{COOH}$$

Thyroxine L(—)-Cysteine, C₃H₁O₂NS (β-thiol-α-aminopropionic acid).

$$\begin{array}{c} \operatorname{NH_2} \\ | \\ \operatorname{HS--CH_2--C-COOH} \\ | \\ \operatorname{H} \end{array}$$

This amino acid is recognized as being present as such in the intact protein molecule but it is not ordinarily obtained as one of the products

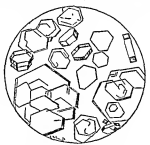


Fig. 45. Cistine.

of protein hydrolysis unless precautions are taken against oxidation, since it is readily converted by oxidation of the —SH group into various sulinic and sulfonic acids and into cystine. The relation between cysteine and cystine may be illustrated as follows:

This reversible oxidation-reduction reaction involving the sulfhydryl group (—SII) appears to have considerable physiological significance. Thus it accounts for the oxidation and reduction reactions of glutathione,

which contains cysteine, and it appears that adjacent polypeptide chains in the protein molecule are linked together through the oxidation of cysteine sulflydryl groups Experiments on cysteine have been grouped with those on cystine

L(--)-Cystine, C₆H₁₂O₄N₂S₂ (di(β-thiol-α-aminopropionic acid))

Cystine is obtained in greatest amount as a product of the hydrolysis of keratin containing tissue such as horn, hoof, feather, and hair It crystallizes in characteristic hexagonal plates which are only very slightly soluble in cold water and in alcohol. It dissolves readily in mineral acids or in alkalis but it is insoluble in acetic acid.

the protein. The greater part of the hydrochloric acid is removed by distilling in vacuo at a temperature between 60° tn 70° C, and the original volume of the solution is restored by the addition of water. A thick aqueous suspension of commercial finishing lime is now sinwly added, care being taken to avoid any considerable rise in temperature, until the mixture has acquired a chocolate color. It is then filtered by suction through a Buchner funnel and the residue washed a number of times with distilled water. The filtrate should be clear and possess a light brown color. Hydrochloric acid is now added to partially neutralize the alkaline solution and it is finally acidified by addition of acetle acid. On standing over night in the icehox, sedimentation of the crude cystine takes place. This is filtered off and le dissolved in a minimum quantity of 5 per cent HCl. The solution is decolorized by boiling for several minutes with a small quantity of charcoal which has been previously hoiled with HCl to remove the calcium phosphate, and the cystine is precipitated by the addition of sodium acetate to the hot solution until a drop of the solution ceases to turn Congo-red paper blue. The mixture is filtered at once and the cystine is washed a number of times with hot water to completely remove the last traces of tyrosine. Typical hexagonal plates of cystlne are obtained. (See Flg. 45.)

Tests for Cystine and Cysteine.

(a) SLLLBAN'S TESTS

Cystine To 5 ml. of the solution under test (containing not more than 0.04 per cent of cystine in approximately 0.1 N hydrochloric acid, at a temperature of about 25° C.) add 1 or 2 ml. of freshly made 5 per cent aqueous solution of sodium cyanide. Mix and let stand 10 mlnutes. Then add 1 ml. of a freshly prepared 0.5 per cent solution of 1,2-naphthoquinone-4-sodium sulfonate, sodium sulfite, etc., as given below for cysteine.

Cysteine. To 5 ml. of solution containing not more than 0.04 per cent of cysteine in 0.1 N hydrociloric acid, add 1 ml. of 1 per cent sodium cyanide in 0.8 N sodium hydroxide. Mix and add 1 ml. of n freshly prepared 0.5 per cent aqueous solution of 1,2-naphthoquinone-4-sodium sulfonete. Mix and add 5 ml. of 10 to 20 per cent solution of anhydrous sodium sulfite in 0.5 N sodium hydroxide. Mix and let stand 30 minutes. A reddish-brown color appears. Then add 1 ml. of a 2 per cent solution of solium hyposulfite (Na.5, O.) in 0.5 N sodium hydroxide. The brown-red color in the presence of cysteine (or cystine treated with sodium cyanide) is converted to a purer red.*

b) Tiers for Sulmittumi. (Sil) Grove. On the addition of a dilute solution of Il, an indigo-blue color appears and disappears almost immediately, d a dilute solution of CuSO., whereupon a transitory violet color appears. lest I or 2 ml. of cysteine solution with a dilute solution of sodium nitroisside and a drop of NaOil. A deep purple-violet color appears but graduy fades after a few minutes.

Add a few drops of a 10 per cent aqueous solution of lead acctate to the vilne (or c) stelne) solution, then render the solution strongly alkaline with per cent NaOli or KOll. Boil for a few minutes. If cystine or cysteine are esent, the solution becomes brown and a black precipitate of PbS appears ad-blackening test).

The reaction requires a high final pH. In case of by droly rates of foodstuffs it is necesto add 1 or 2 ml. of 5 N sodium by droude just before adding the final reducing agent, from by possible (Nais-O.). As a rule it is systime that is found in hydroly rates.

L(-)-Methionine, $C_2H_{11}O_2NS$ (γ -methylthiol- α -amino-n-butyric acid).

When methionine is treated with concentrated sulfuric acid, the methyl group is split off and the amino acid homocy steine is ultimately obtained

Homocysteine is similar to cysteine in many ways, and its formation and quantitative determination may be used for the quantitative determinution of methionine While homocysteine has not as yet been isolated from proteins, there is good evidence that it is an intermediate in the biological transformation of methionine into cystine (see p 1030)

EXPERIMENT ON METHIONINE

Methionine can be detected in a mixture of amino acids by the following method (Bolling a modification of McCarthy-Sullnan) To 75 ml of unknown, add the following reagents in order, mixing after each addition 15 ml of 5 N NaOli, 15 ml of 1 per cent glycine, and 0 3 ml of 10 per cent sodium nitroprusside (freshly prepared) Place the tube in a water bath at 37' to 40° C. for 15 minutes, cool in ice water for 2 to 7 minutes, and add 3 ml of 6 % HCl Shake for I minute and let stand at room temperature for 15 min utes A reddish purple color indicates methionine Tryptophan Interferes.

L(-)-Tryptophan, C11H12O2N2 (β-3-indole-α-aminopropionic acid).

small amount of glyoxylic acid added, and sulfuric acid then stratified on the bottom of the tube, a reddish-violet ring will appear at the juncture of the two liquids. This is also known as the glyoxylic acid test (see p. 170). Pure tryptophan will not give this test except in the presence of a trace of ferric or curre ions.

(b) ALDEHYDE REACTION (VOISENET-RHOOS) Tryptophan will also give color reactions in the presence of aromatic aldehydes. With p-dimethylaminobenzaldehyde in sulfuric acid it gives a red-violet color. These color reactions are apparently due to the presence of the indolering in tryptophan. The indole



FIO 46 TRYPTOPHAN
From Keenan J Biol Chem 62 163 (1924)

ring, being a combination of the benzene and pyrrole rings, probably owes its clin being properties to the latter ring. For other tryptophan reactions see Chapter 16

L(-)-Proline, C,H,O,N (pyrrolidine-2-carboxylic acid).

Proline is easily soluble in alcohol and in cold water. It has a sweet taste and melts at 153° to 154°

L(-)-Hydroxyproline G₆H₉O₄N (4-hydroxy-pyrrolidine-2-corbox) lic ocid).

It is very difficult to separate hydroxyproline from the other amino acids. It is easily soluble in water and soluble with difficulty in absolute alcohol.

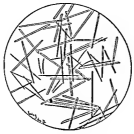


FIG 47. PROLINE

L(-)-Aspartic acid, C_iH₁O₄N (α-aminosuccinic acid).

COOH CH: H—C—NH:

This amino acid, unlike the others thus far considered, is strongly acidic on account of the predominance of the carboxyl group. The chief source



Fig 48 ASPARTIC ACID

of this acid is the monoamide, asparagine, which is very widely distributed in the vegetable world, being particularly abundant in the asparagus plant and in lentil sprouts. It is more than likely that the amide (aspara-

gine) rather than aspartie acid occurs in the protein molecule, but is quickly hydrolyzed into aspartie acid and ammonia during the process of protein cleavage.

L(+)-Glutamic acid, CsH2O4N (α-aminoglutaric acid).

This acid, when obtained after the hydrolysis of the protein molecule, is largely a secondary product. The primary constituent of the protein molecule is undoubtedly glutamine, the amide of glutamic acid, which accounts for the greatest part of the total glutamic acid found after bydrolysis of the protein. Glutamic acid is present in practically all proteins,

usually in fairly large amounts, being present to the extent of 47 per cent in gliadm, a protem found in wheat, and to the extent of 23 3 per cent m casein. The sodium salt of glutamic acid is widely used commercially for flavoring soups, sances, and food concentrates.

Glutamic acid is found combined with glycine and cysteine in the

glutathione molecule (see p 321) Glutathione has been shown to be a tripeptide having the following structure

Attention is called to the fact that the glutamic acid is linked to the amino group of cysteine through its γ carboxyl group



L(-)-Histidine, $C_6H_9O_2N_3$ (β -imidazole- α -aminopropionic acid).

$$\begin{array}{cccc} & & & & & & \\ & & & & & & \\ CH = C - CH_2 - C - COOH \\ NH & N & H \\ & & CH \end{array}$$

Histidine is found in small quantities in practically all proteins, but is present to the extent of 8 per cent in hemoglobin. It combines with a variety of substances, forming, for example, mono- and dihydrochlorides, also compounds or double salts with platinum chloride or silver nitrate which are particularly valuable for obtaining a pure crystalline com-

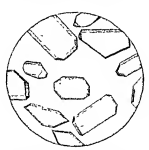


Fig. 50. HISTIDINE DIHYDROCHLORIDE.

pound. It also forms characteristic compounds with pieric acid, phosphotungstic acid, nitrobenzoyl chloride, 2,5-dichlorobenzene sulfonic acid, nitranilic acid, etc.

L(+)-Arginine, C_δH₁₄O₂N₄ (δ-guanidino-α-aminovaleric acid).

Arginine is present in all tissuo proteins. It is hydrolyzed into urea and the amino acid ornithine (a,3-diaminovaleric acid) by the enzyme arginase which is found in the hyer.

This reaction is thought to be of major importance in connection with the formation of urea by the body (see Chapter 33 for a discussion of this and other metabolic relationships of arginine).

EXPERIMENTS ON ARGININE

1. Itolation of Arginine Flavianate (Kossel). Hydrolyze a 25-g, portion of gelatin by boiling with 250 mil. of 18 per cent hydrochloric acid under reflux for 18 hours. Remove the excess acid by repeated concentration in vacuo. Take up the residue in 250 ml. of hot water and decolorize with 5 g, of charcoal. Bring the filtrate to a volume of 250 ml. and add a saturated aqueous solution containing 200 g, of flatianic scid (2,4-dintro-1-naphthol-7-sulfonic acid) at room temperature. Allow the precipitate to form in the cold for 5 days, stirring from time to time. Fiter off the yellow precipitate and wash with a little cold water. Dissolve the washed precipitate in hot water with the aid of a minimal quantity of 4 per cent ammonia. While the solution is still bet adds sufficient 29 per cent hydrochloric acid to neutralize all the ammonia. Arginine flavianate crystallizes in shining yellow plates from the hot solution.

L(+)-Lysine, C_εH₁₄O₂N₂ (α-ε-diaminocaproic acid).

Lyane is one of the basic aminn acids, possessing a predominance of amino groups over acidic or earboxyl groups. Histidine, arginine, and hisne have been called "the become basis." Lyane is present in most proteins of animal origin. It is notably absent from zein and present in rather small amounts in gladin.



FIG. 51, LYSINE PICRATE.

Gitrulline, $G_6H_{13}O_3N_3$ (δ -carbamino- α -aminovaleric acid).

$$\begin{array}{c} NH_{2} \\ NH_{2}-C-NH-CH_{2}-CH_{7}-CH_{2}-C-C00H \\ 0 \\ \end{array}$$

Citrulline was obtained by Wada in 1930 from watermelon juice. Little is known concerning its distribution in proteins Free citrulline is present in small amounts in liver and in blood. Interest in citrulline at the present time is based largely upon its relation to arginine and the processes of urea formation in the animal body (see Chapter 33).

Newer Amino Acids. The application of newer research techniques such as paper chromatography for the isolation and identification of amino acids in hydrolyzates, and the use of microorganisms as nutritional test subjects, have led to the discovery of a number of amino acids and related compounds not hitherto identified among the products of protein hydrolysis. Among those of biological interest, mention may be made of:

α-Aminobutyric acld found in the brain and central nervous system, where it is formed by decarboxylation of glutamic acid. 10 It has also been shown to be produced enzymatically from pi-threonine. 11

Lanthionlne (THIODIALANINE) from wool;12

Cystathlonine, a condensation product of homocysteine and scrine intermediate in the synthesis of cysteine;13

Ergothloneine, found in ergot, blood (1.8-1.95 mg. per 100 ml. in human blood),¹⁴ and in the seminal plasma of the boar;¹⁵

¹⁰ Roberts and Frankel. J. Biol. Chem., 187, 55 (1950); 188, 789 (1951); 190, 505 (1951).

¹¹ Lien and Greenberg. J. Biol Chem. 195, 637 (1952); 200, 367 (1953).

Horn, Jones, and Ringel: J Biol. Chem., 138, 141 (1941).
 Binkley, Anslow, and du Vigneaud: J. Biol. Chem., 143, 559 (1942).

¹⁴ Hunter. Brochem. J., 48, 265 (1951).

¹ Mann and Leone: Brochem. J., 53, 140 (1953).

3,5,3 Truodothyronine from thyroid glands 16

Methionine sulfoximine, the toxic factor isolated from flour matured by treatment with introgen trichloride 17

Tabtoximine (a e Diamino & hydroxypimelic acid) a phytotoxic compound found in certain discased tobacco plants 15

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* Woolley B haff er and B aun J B of Chem 198 807 (13-2)

Proteins: Their Structure and General Reactions

The results of investigations on the hydrolysis of proteins indicate that the protein molecule is composed almost evelusively, if not entirely, of α -amino acids. Investigations of this type, which involve the complete tearing down of the protein molecule, yield the constituent amino acids but throw practically no light on the question as to the order in which these acids are linked together to form the intact protein moleculo with its characteristic elemical, physicochemical, colloidal, and biological properties. In spite of this defect these researches are of the highest importance since they yield the actual "building stones" of the protein molecule

In recent years emphasis has been placed on the study of the physicochemical behavior of the intact protein. The difficulty involved in the formulation of an acceptable theory for the structure of the protein molecule becomes apparent when we list the great number and variety of properties and reactions of proteins that such a theory is called upon to oxplain Of these, we may mention especially the following (1) The hydrolysis of proteins by acids, alkalies, and proteolytic enzymes into their constituent amino acids, (2) the small proportions of free amino and carboxyl groups in the intact molecule, (3) the large increase in both amino and carboxyl groups that accompanies the hydrolysis of the protem, (4) the combinations of proteins with acids, bases, and many other classes of compounds, (5) the large size and colloidal nature of the protein molecule, together with the complex colloidal behavior of protein solutions, (6) the complex solubility relationships of the various classes of proteins, (7) the sensitiveness of proteins to chemical and physical agents such as acids, alkahes, alcohol, heat, mechanical shaking, ultraviolet light, etc., (8) the immunological reactions of the proteins, and (9) their highly specific enzymatic, hormonal, and other biological properties. In the following sections the modern conceptions of protein structure will be discussed briefly together with the more important reactions and properties of proteins which make them such vital constituents of hving protoplasm

Structure of Protein Molecule: Peptide Linkage. It was pointed out by Hofmerster, in 1902, that there are three concerable types of hinkago by which individual amino acids might be joined together in the protein molecule

The first type of hukage involves direct union between earhon atoms.

This type of union is very unlikely since bonds between carbon atoms are not attacked by protoclytic enzymes which hydrolyze native proteins It is difficult, also to understand how a molecule having such a structure could be broken down by hydrolytic agents into such definite structural units as the polypeptides and amino acids

The second type of linkage that suggests itself is a linkage of carbon atoms by means of an oxygen atom, as in the ethers, esters, and anhy drides

This type of union is also improbable since, in view of the number of its carboxyl groups, the protein molecule does not contain sufficient oxygen to account for a major linkage of that kind Moreover, since such a link age does not involve the amino groups, the latter should be much more abundant in the intact protein molecule than they actually are

this leaves, as the final possibility, the linkage of carbon atoms by means of a nitrogen atom

Of the various possibilities for such a linkage, that resulting from the condensation of the amino group of one acid with the carboxyl group of another, which Fischer named the peptude linlage, is the only one which is in accord with the experimental facts. Fischer devised several ingenious methods for condensing amino acids in this manner and prepared a large number of di-, tri , and polypeptides some of them containing as man) as 18 molecules of amino acids Many of the synthetic polypeptides prepared by Fischer, Bergmann, and others are identical with polypeptides isolated from partially hydrolyzed proteins

According to modern conceptions of protein structure the peptide bond is the predominant bond in the protein molecule. This point of view is based on a great mass of experimental evidence, of which only a brief review may be given here The structural formula of the tetrapeptide given below indicates clearly that only those amino and earhoxyl groups which are at the ends of chains are free When the molecule contains diamino or dicarboxylic acids the additional amino or earboxyl groups

Glycyt

Tyrosine

either may remain free or may be the starting points for side chains of various kinds. Analyses of proteins indicate that the number of free amino aud carhoxyl groups in the intact molecule, determined by Van Slyke's nitrous acid and Sørensen's formol titration methods, is substantially what would be expected of a molecule built up of amino acids joined together hy means of peptide linkages. Furthermore, hydrolysis of proteins hy acids or enzymes results in the liheration of equal numbers of amino and carboxyl groups, such as would arise during the hydrolysis of peptide bonds. The hiuret reaction, which is characteristic of proteins and some of their decomposition products, is given by many of the synthetic polypeptides and practically disappears when all these substances are completely hydrolyzed.

Relation of Structure to Properties of Proteins. The generally accepted theory of Hofmeister and Fischer that proteins consist of chains of amino acids joined to each other through their amiuo and earhoxyl groups does not hy itself seem to explain the unique chemical, physical, and hiological differences among proteins in nature. Why are keratins so resistant to dilute acids and proteolytic enzymes? How does the muscle protein, myosin, function in the contraction of muscle? What is responsible for the immunological specificity of proteins, for the enzyme action of eatalase and pepsin, for the toxicity of tohacco mosaic virus, and so forth? The hypotheses which are mentioned hriefly below are attempts to an-

swer some of these questions.

A. PROTAMINE NUCLEUS HYPOTHESIS OF KOSSEL AND SIEGFRIED. These investigators helieved that all proteins were huilt around a nucleus of the three amino acids arginine, histidine, and lysine, and that arginine was the most important member of this triad. Block has shown that one group of proteins, eukeratins, can he characterized by the relative constancy of the ratios of arginine to lysine to histidine. The location of these three amiuo acids or their mode of action in the formation of the eukeratins remains unknown. It should also he pointed out that no protein devoid of arginine has as yet been reported, although some large peptides are lacking in this amino acid.

B. DIKETOPIPERAZINE HYPOTNESIS OF ABRERHALDEN. Amino acid anhydrides (diketopiperazines, see p. 130) can be obtained from proteins under certain conditions. These compounds are readily formed by condensation of two amino acids or by eyelizing a dipeptide. Abderhalden suggested that proteins are composed of diketopiperazine-containing units which are held together by secondary valences. This structure was intended to account for certain properties of proteins, especially the hydrolysis to peptones by pepsin. Unfortunately for this theory, amino acid anhydrides are not hydrolyzed hy proteolytic enzymes.

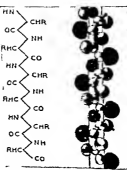
C. STEREOCHEMICAL ORGANIZATION OF THE PROTEIN MOLECULE (FOA, MEYER AND MARK, ASTBURY, NEURATH, WHINCH, PAULING). Tho use of the x-ray to study the fine structure of proteins, especially in the hands of W. T. Astbury, has given us a much clearer insight into the organization of the peptide chains in the protein molecule. A short section of such a chain in the fully extended position is shown in Fig. 52. In general these peptide chains are pictured as either fully extended (silk fibroin, or other 

FIG 52 SCALED MODEL OF A FULLY EXTENDED POLYPEPTIDE CHAIN VIEWED



account for these distinctive v-ray patterns, all of which suffer from the disadvantage of not meeting all the experimental evidence In 1951, Linus Pauling and his co-workers, Corey and Branson, employing as sole restrictions to their formulation the accepted interatomic distances and covalency angles, the planarity of the amide groups, and the linearity of the O-hydrogen bond and assuming that --N--H the maximum possible number of intramolecular hydrogen bonds would be formed, arrived at certain helical structures 1 One of these gave 3.7 amino acid residues per complete helical turn. This resulted in 13-membered rings, closed by -CO- and -NHgroups in hydrogen-bond linkage as shown in perspective and in plan (Figs 53 and 54, respectively) This so called α-lielix has been rather definitely established to occur in hair, horn, and other proteins of the a-keratin class (including muscle) as well as in hemoglobin and many other globular proteins such as serum alhumin, insulin, lysozyme, and chymotrypsm Another proposed helical formation, the y- or 51-residue helix, is now believed to be too unstable to exist in proteins How-

appears to be characteristic of silk and stretched hair. It is generally helieved that the gamma globulin of blood is able to assume different shapes under the impact of foreign antigens while retaining its amino acid composition and arrangement unmodified. It seemed that too rigid specification of the peptide (backhone) coiling of gamma globulin would be incompatible with its known lability with respect to the formation of antibodies. The 3.7 amino-acids residue helix seems capable of overcoming this difficulty since it may be transposed into many different patterns by starting the helix with the same amino acid but at geometri-

ever, an "antiparallel-chain pleated sheet" structure1a



Fig 54. PLAN OF THE 37-RESIDUE HELIX.

^{&#}x27;10 53 Thie Hetx wirit 3 7 RestDULS 1ER TURY

1 Pauling Corey, and Branson Proc Natl Acad Sci. 37, 207
(1931) and personal communication
1 Pauling and Corey Proc Natl lead Sci. 37, 729 (1931)

cally different points. To visualize the 3.7 helps better, it may be compared to a spiral staircase with each amino acid residue as a step. The height of each step is 1.5 $\mathring{\Lambda}$ and the height of each turn is 5.4 $\mathring{\Lambda}$ making 3 6 or 3.7 steps per turn. It thus takes 18 steps or 5 turns for a step to be found

exactly in a vertical line above the starting point.

The closely knit structure of highly organized soluble globular proteins is broken down by denaturing agents such as strong sodium hydroxide. The peptide chains then become a mass of disorganized fibrils in solution. If this alkaline protein solution is passed through a small capillary, these fibrils orient themselves with their long axes parallel to the direction of flow. If the alkaline solution is then extruded through a small orifice into an acid or other coagulating hath, the protein fibrils unite with each other parallel to their long axes and a typical macroscopic thread is formed. This is the essence of the production of synthetic fiber, "wool," etc., from soluble globular proteins such as casein and soybean.

The three main types of cross linkage in proteins appear to be (1) Dithio bridges formed by cystine, (2) Salt bridges formed by the carboxl groups of aspartic or glutamic acid and by amino groups of lysne or granido groups of arginine, and (3) Hydrogen bonds between peptide link-

ages and between polar groups of the amino acids:

From this presentation it is apparent that the solution to the problems of protein structure and function has not yet been found, but that the Hofmenstr-Fischer peptide theory, coupled with the known electrical and chemical properties of the peptide linkage (—NH—CO—CHR—) and of the specifie side chains of the amino acids, offers a reasonable basis to account for many of the properties of natural and of isolated proteins.

Proteolytic Enzymes. The most convincing evidence that the peptide linkage is the principal type in the protein molecule comes from the fact that enzymes are able to hydrolyze synthetic polypeptides of known structure, liberating equal amounts of amino and carboxyl groups, just as they do when acting on native proteins. Our knowledge on the structural specificity of enzymes is due in large part to the brilliant researches of Bergmann and his co-workers. The table given below illustrates the specificity of enzymes.

ACTION OF PROTECLYTIC ENZYMES (BERGMANN)

Enzyme	Requisite Peptide Chain	Requisite Amino Acid		
Pepsin	CONIL—CH—CO NII—	Tyrosine or Phenylalanine		
Cathepsin A(I)	-CONII-CH-COOH + H ₂ N-			
Trypsin Cathepsin B(II) Papain H-S	Same as above except for R group	Lysine er Arginine		
Leucyl amino- peptidase (Cathepsin III)	R H ₂ N-CH-CO'NH- HO'H R H ₂ N-CH-COOH + H ₂ N-	Leucine		
Carboxypeptidase (Cathepsin IV)	R -CO.NH-CH-COOH HO.H HO.H R -COOH + Han-CH-COOH	Tyrosine or Phenylalanine		

Up to the time of these investigations it was believed that the size and not the amino acid composition of the protein or polypeptide was the controlling factor in enzymatic hreakdown. Thus pepsin was thought to act on proteins of high molecular weight such as fihrin, caseiu, etc., but not on the smaller protamines, while trypsin split the smaller molecules, such as the protamines, and crepsin hydrolyzed peptides of still lower molecular weight. It is now evident that pepsin fails to hydrolyze protamines not because of their molecular size but hecause these proteins are

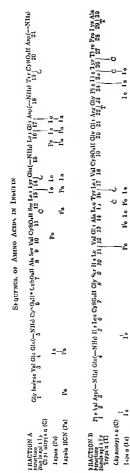
deficient in tyrosine and phenylalanine while the ample supply of arginine and lysine in protamines permits the action of trypsin Likewise a mixture of polypeptides containing a relatively high proportion of free carboxl groups would be split by carboxypeptidases (erepsin?) such as the aromatic carbox peptida e shown in the table above In each case, the ratio of carbox il to amino groups liberated is unit; The resistance of certain proteins such as silk fibroin hair keratin, etc to digestion by proteolytic enzymes is not explainable on the hasis of amino acid composition but is due to the close packing of the protein molecules which then offer rela tively little surface to the enzyme If the molecular structure is disor ganized by mechanical or chemical means these proteins are readily by drolyzed by the proteolytic enzymes

In the current hypothesis of enzyme action it is assumed that the en zyme substrate complex is formed by the honding of complementary sur facts Thus the mechanism would depend primarily on the steric arrangement of the molecular groups of the substrate their shape and mutual positions and only secondarily on the chemical nature of the bond under

c ang livdrolvsis

The Composition of Insulin In spito of all the chemical and physicoclemical studies on many proteins only one protein has been so thor outhly investigated that its amino acid sequence is completely known In a series of brilliant researches Sanger and his co-workers at Cambridge Lucland have elucidated the chemical structure of pure crystalhae in sulm In brief they broke the -SS-- honds which hold the peptides of moulin by oxidation with performic acid This procedure not only hreaks the -SS- bonds but also oxidizes the eysteine residues to cysteic acid residues. The resulting oxidized insulin was then separated into two frac tions fraction 1 contained only glycine at the amino end of the chain fraction B contained only phenylalanine as the \ terminal amino acid these two fractions were then subjected to partial hydrolysis hy dilute acid or by enzymes When the hydroly tic products (amino acids and small peptides) were separated and identified by means of paper chromatog raphy Sanger Thompson and Tuppy were able to fit these together to give the exact sequence of all the amino acids in each of the peptides Their results are given in the accompanying table

Molecular Weights of Proteins Many of the properties of protein solutions especially those connected with their colloidal behavior are intimately related to the size and molecular weight of the protein The determination of the molecular weights of the proteins involves man't difficulties owing to the complex solubilities of the proteins and the collor dal nature of their solutions Thus for example the ordinary physico chemical methods such as the raising of the boiling point or lowering of the freezing point are either mapplicable or yield misleading results when applied to proteins because of the effects of traces of salts and other im purities on determinations made by those methods. However, the minimal melecular weight can be of tained from chemical especially amino acid at altace By assuming that the protein inolecule contains of e atom of an e cincut or one me ecule of the amino acid present in least amount



* The cysteic and exists in the insulin molecule as cystine indicates offier bonds split by enzymes indicates major sites of enzyme action

Structure Bonds spi t ly Trype n (T) RACTION B

1 . (10)

I cpen (Pe)

the minimal molecular weight may be calculated from the following relationship

Minimal molecular weight of protein

= 100 × Atomic weight of element
Percentage of element

= $100 \times \frac{\text{Molecular weight of amino acid}}{\text{Percentage of amino acid in protein}}$

On the other hand the relatively large particle size of most proteins has permitted the development of methods for estimating their size and molec ular weight which could not be used for the average organic compound These are

 Molceular weight from osmotic pressure (Sørensen, Roche, Adar, Greenberg, et al)

2 Molceular weight from sedimentation equilibrium and from sedi mentation rate and diffusion (Svedberg, Peterson)

3 Molecular weight from diffusion and viscosity (Northrop and Anson, Neurath, et al)

4 Molecular size and shape from dielectric constant dispersion curves (Williams, Oneley) 5 Molecular size from x ray diffraction data (Crowfoot, Fankuchen)

The osmotic-pressure method yields the mean molecular weight when carned out under suitable experimental conditions This procedure, which ean he earned out with ordinary laboratory equipment, suffers from the disadvantages that it gives no indication of homogeneity of the substance investigated and that it becomes increasingly insensitive with increasing molecular weights

Every molecule in solution is subject to thermal forces which result in diffusion of the substance away from a center of high concentration toward one of lower concentration In addition to these thermal forces the molecule is subject to gravitational forces which tend to cause the molecules to sediment It is thus apparent that whether a substance will remain in solution or not is dependent upon the relative strengths of the gravitational (settling) and thermal diffusion forces By increasing the speeds attainable in the ordinary laboratory centrifuge, Svedberg, Beams, and others have constructed very high speed contribuges (ultracentrifuges) which are able either just to halance the diffusion forces of a protein molecule in solution (sedimentation equilibrium) or actually overcome these thermal forces and cause the protein to sediment at a definite rate (scdimentation velocity)

If the concentration diffusion constant, and approximate shape of the protein are known the molecular weight or particle size of the protein or other colloid can be calculated either from sedimentation-equilibrium or sedimentation-velocity data (see Fig 55) One significant contribution of this method is that it is capable of separating particles of different sizes and weights and thus indicating whether the protein under investigation is mono- or polydisperse Many proteins which were previously con sidered to be monodisperse are now known to be mixtures of products varying widely in particle size. It should be recalled, however, that protein particles of the same size (monodisperse) may differ in amino acid composition, thus making the protein a mixture; on the other hand protein particles of different sizes (polydisperse) may have the same amino acid composition and thus be chemically pure. Analogies exist in organic chemistry; thus leucine admixed with a little isoleucine would be monodisperse, although impure; whereas acetic acid dissolved in benzene would be both monomeric and dimeric acetic acid (polydisperse) and yet be all the same compound.

Molecular weights of proteins have been enleulated from diffusion and viscosity data. Both diffusion and viscosity are functions of the size,

shape, and degree of hydration of the molecule.



Fig. 55, Schimentation in the Ultragentrifuge of a Monodisperse Sol (Hemoeyanin) and a Polydisperse Sol (Gold),

From Svedberg' Colloid Chemistry, 2nd ed Reinhold Publishing Corp., New York, 1928.

Proteins, among other substances, can be oriented in an electric field. If the direction of the field is reversed the molecule will reorient in the opposite direction. As the nate of change of direction of the current is increased (i.e., increasing frequency), more and more of the molecules fail to orient properly and the electrical properties of the solution change. The changes in the dielectric properties of a protein solution are also dependent upon the size, shape, and viscosity of the molecule as well as the alternating frequency. Such data have been used to calculate the molecular weight of some proteins.

The final method mentioned above for the estimation of molecular size is based upon the fact that a single protein crystal, when placed in the beam of an x-ray, will cause the beam to be somewhat deflected. From this, the dimensions for a unit cell are obtained and from the density of the protein its mass is computed. The molecular weight is calculated by dividing this mass by that of the hydrogen atom (1.66×10^{-24}) . The results indicate the molecular weight of tho protein or some multiple thereof.

Additional methods for determining molecular weight and shape of

proteins are molecular shape from flow-birefringence data, molecular weight from light scattering of proteins, and dimensions of proteins from electron micrography

It should be pointed out that values given in the table of molecular weights below may be multiples or fractions of the true molecular weights of the proteins. These methods probably measure the size of the protein particles with considerable accuracy under the particular experimental conditions, but the results do not necessarily indicate the true molecular

MOLECULAR WEIGHTS* OF PROTEINS

	Method					
Protein	Cl em- zcal	Ultra- centrifuge	Osmotic Pressure	Diffu *ton	l secos tlj	\ I ay
Lactalbumin Cytochrome C Vlyoglobin	13 000	17 400 15 600 16 900				
Gliadin Hordein	1 1	27 900 27,500	40 000	27 500		i
Zain Concanavalin B Crotoxin Inaulin† Pepsin Ovallutuin \$ Lactoglobulin Serum globulin (terse) Serum globulin \(\gamma\) Catalasse Lidestin Lrease Nucleohistone calf Tobacco messae virus	23 000 36 000 34 200 40 000 33 400 73 000 66 700 164 000 53 000	153 000 250 000		33 400 71 000 63 000	40 000	36 300 39 500 37 500 34 000 82 000 67 500

Many values are averages from the literature
 See discussion in text.

weight. The discrepancy between apparent molecular weight and true molecular weight is illustrated in the case of insulin. This highly purified protein was believed to have a molecular weight of from 35,000 to 45,000 by the commonly employed physical methods Modifications of these procedures suggested at first a molecular weight of 12,000 and then of 6,000 like latter value agrees most closely with the chemical analyses of insulin which indicate one residue of glycine, alanine, and isoleucine per mole of protein.

¹ Nature 178 518 (1952)

Nature of Protein Solutions, Studies of the behavior of the scrum globulins led Sørensen, some years ago, to conclude that euglobulin and nseudoglobuliu were reversibly combined in a loose chemical combinatiou, E.P., in which E and P represent euglobulin and pseudoglobulin complexes, respectively, combined in the relative proportions of p and q. Subjecting serum to such procedures as dialysis, or fractionation with ammonium sulfate, results in a shifting of the proportions of E and P with the resultant formation of more soluble and less soluble complexes of the two proteins. Sørensen failed, even after repeated fractionations, to prepare samples of either euglobulin nr pseudoglobulin that were completely free from the other protein. These results with the globulins led to a series of investigations of other prateins with the result that bighly purified preparations of scrum albumin, casein, and gliadin were each found to consist of mixtures of an unknown number of proteins of similar character combined in a reversible manner. Such proteins, according to Sgrensen, represent "reversibly dissociable component systems" and may be represented by the formula A.B.C. . . . , in which A, B, C, etc., represent components of a definite character and composition (e.g., polypeptides) while x, y, z, etc., indicate the number of such components in the more complex system. In each component, the atoms or groups of atoms, such as amino acids, are linked together by means of strong chemical honds, whereas the complexes are formed by the union of these components through weak, residual valences. Chemical or physical agents that act ou the chemical bonds produce irreversible changes in the protein molecule whereas the residual valences respond to changes in salt concentration, pH, or temperature in a reversible manner. Although the various fractions obtained by the fractionation methods employed possess the essential properties of the initial material, they exhibit variations in physical properties and chemical composition that are considered to be duo to the varying amounts of the individual components in each fraction. In no case has Sørensen succeeded in isolating a component which could not he further fractionated by appropriate methods.

Applying these conceptions to biological systems, we find that not only may comparatively simple components combine with each other by means of their residual valences to form protein complexes, but these complexes may themselves combine to form still more complicated structures. In serum, for example, we probably have not only such component systems as alhumins and globulins, which may be isolated by suitable methods, but also more complex systems in which these proteins are combined in varying proportions not only with each other but with other serum constituents, such as the lipides. In protoplasm, instead of relatively inert, independent substances, there are probably complex systems composed of protein, lipide, and carbohydrate in equilibrium with each other (orosins) and constantly shifting in response to changes in environment. The multiplicity and flexibility of such systems may he of profound importance in determining the adaptability of the organism to its environment.

If we accept Sørensen's views on proteins, we can understand the extraordinary difficulties encountered in the isolation of individual proteins from such complex materials as serum and egg white, and can explain results obtained in Svedberg's laboratory on these substances Svedberg found, for example that there was no substance in fresh egg white with a molecular weight corresponding to that of the crystallized oralbumin, but that this substance appeared only after the egg white was treated with ammonium sulfate as in the erystallization method employed Similarly Svedberg found that half saturation of serum with ammonium sulfate precipitated a globulin fraction which was homogeneous and had a defi nite molecular weight and that the euglobulin and pseudoglobulin appeared only after this substance was subjected to further fractionation processes. It thus appears that the proteins with which we are familiar exist in nature only as part of more complex systems, and that even these purified proteins may themselves be complexes formed by the union of several simpler components. The problem of determining the structure and properties of the protein molecule thus depends upon the development of suitable methods for the isolation, in pure form, of the compara tively simple components

The complexity of the serum proteins is illustrated by the large num ber of components which one is able to obtain by suitable fractional precipitation of serum. This subject is discussed in detail in Chapter 22

Colloidal Behavior of Protein Solutions The proteins form col loidal solutions of the type known as emulsoids or hydrophilic colloids One of the most characteristic properties of emulsoids as pointed out in Chapter 1, is that such systems have two stability factors charge and hydration, either of which is capable of keeping the particle in solution Individual proteins show marked differences in the hydration of their particles Most proteins are soluble in dilute acids and alkalies the par ticles acquiring positive or negative charges depending upon the pH of the solution For every protein there is a definite characteristic pH known as the isoelectric point, at which the particles are electrically neu tral and cease to migrate to the poles of an electrical field. Although all proteins are least soluble at their isoelectric points certain proteins such as gelatin or ovalbumin remain in solution when brought to their respec tive isoclectric points. On the other hand less soluble proteins such as casein and edestin remain in solution only at acid or alkaline reaction and precipitate when their solutions are brought to the isoelectric point Ovalbumin thus behaves as a typical emulsoid forming stable solutions of neutral particles Casein acts more like a suspensoid the particles of which flocculate when their charges are neutralized. The effect of hydra tion and charge on the colloidal properties and stability of protein solu tions is illustrated in Fig 56 adapted from Kruyt

The determination of the true nature of protein solutions is complicated by the fact that the proteins are unique colloids since in addition to their spically colloidal properties they function as amphotoric electroly to because of the free amino and carboxyl groups which their molecules contain \ \text{gr.at} \ \text{deal} \ \ \text{of confusion custed as to whether the reactions of proteins with acids and based are the mild reactions taking place in stoichiometric proportions or colloidal reactions following various laws of adsorption \ \text{Vuch of this confusion is probably due to differences in the

nature of the particles present m different protein solutions. The work of Pauli, of Sørensen and of Loeb, and more especially the fine work of Svedbeig, discussed above, indicates that in sofutions of such proteins as ovalbumin, hemoglobin, and edestin, the material is dispersed in the form of individual molecules of protein and not as molecular aggregates. Such proteins therefore form molecular solutions whose colloidal properties are due entirely to the companatively large size of the individual molecules. Solutions of other proteins, such as gelatin and casein, probably contain molecular aggregates as well as individual molecules. This point of twe is supported by the growing mass of evidence obtained from

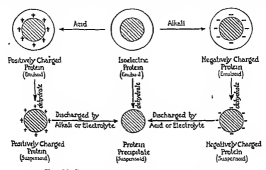


FIG 56 COLLOIDAL BEHAVIOR OF PROTEIN SOLUTIONS

ultracentrifugal and ultrafiltration experiments as well as from numerous studies on the physicochemical properties of protein so utions

Behavior of Proteins as Amphoteric Electrolytes: Isoelectric Points of Proteins. It is currently behaved that proteins behave as molecular solutions of amphotene electrolytes which exhibit typical colloidal properties because of the large size of the individual molecules These large molecules contain reactive amino and carboxyl groups which are capable of entering into true chemical combination with acids and bases In choosing between the chemical and colloidal interpretations of protein reactions, it is well to consider, as pointed out by Svedberg, that colloidal and chemical forces are fundamentally the same, since both are electrical in character and depend upon the attraction and repulsion of negative electrons and positive nuclei. It is quite probable that different proteins vary in their reactions because of the fact that solutions of certain proteins undoubtedly contun individual inolecules, but others contain molecular aggregates of various sizes. In the present state of our knowledge of protein chemistry we may assume that the proteins exhibit both chemical and colloidal properties, the former being due to the presence of reactive groups in the protein molecule and the latter being dependent upon changes in the charge and hydration of the particles in solution, whether molecules or molecular aggregates

Considered as amphoteric electrolytes, the proteins combine with acids and bases to form salts which ionize into a colloidal protein ion and one or more crystalloidal ions. According to the work of Loeb, Sørensen, Pauli, Michaelis, Cannan, and their co-workers, this combination takes place in definite proportions in accordance with the laws of classical elemistry A protein at its isoclectric point is either entirely without charge—1e completely un ionized—or else, according to the zwiterion hypothesis of Bjerrum, carries equal numbers of positive and negative charges due to complete dissociation of equal numbers of acid and basic groups in the molecule. Isoclectric protein may thus be represented by the formulas

Here R stands for the nucleus of the protein molecule. The addition of acid to isoelectric protein results in the formation of a protein salt (e.g., protein chloride) which ionizes into a positive protein ion and a negative

$$\begin{array}{ccc}
\text{COOH} & \text{COOH} \\
R & + \text{HCl} \rightarrow R \\
& \text{NH}_{2} & \text{H}
\end{array}
= \begin{bmatrix}
\text{COOH} \\
R & \\
& \text{H}_{3}
\end{bmatrix}^{+} + \text{Cl}^{-}$$

acid ion. In the same way the addition of alkali to isoelectric protein results in the formation of a salt which ionizes into a positive metal ion and

$$\begin{array}{c} \text{COOH} \\ \text{R} \\ + \text{NaOH} \rightarrow \text{R} \\ \\ \text{NH}_2 \end{array} \rightleftharpoons \begin{bmatrix} \text{COO} \\ \text{R} \\ \\ \text{NH}_2 \end{bmatrix} + \text{Na}^+ \end{array}$$

a negative protein ion. In solutions acid to their isoelectric points therefore, the proteins exist as positively charged ions, capable of combining with negative ions to form salts while in solutions alkaline to their isoelectric points proteins exist as negatively charged ions which can combine only with positive ions. Loeb added solutions of silver nitrate and potassium ferrocyanide to powdered gelatin which had been previously brought to various hydrogen ion concentrations by soaking in appropriate solutions of acid and alkali He showed that gelatin combines with silver only when the solutions are on the alkaline sine of its isoelectric

^{*} The above reactions make use of the older formula for isoelectric protein. The same products are formed according to the zwitterion hypothesis the only difference being in the meet anism involved (See section on anismo acids Chapter 4).

point (pII > 4.7), and with ferrocyanide only on the acid side of its isoelectric point (pII < 4.7). At the isoelectric point gelatin behaves as though it were un-ionized, since it does not combine with either positive ions (cations) or negative ions (anions).

The isoelectric points of the proteins are of especial significance in protein chemistry, because the properties of the proteins undergo unique changes at these points. It has already been pointed out that proteins are ionized and can enter into chemical combination only in solutions which are acid or alkaline with respect to their isoelectric points. The solubility of the proteins, especially those proteins which resemble the suspensoids in their colloidal behavior, is either negligible or at a minimum at the isoelectric points. Other physical properties of the proteins such as viscosity, osmotic pressure, swelling, etc., are also at a minimum at the isoelectric points. According to Loeb these properties are dependent upon a Donnan equilibrium set up between the particles and the surrounding dispersion medium. The isoelectric points of a number of the more common proteins are given in the following table.

ISOLLICING POINTS OF SOME COMMON PROTEINS

Protein	pII		
Ovalbumiz	1 55-4 90		
Edestin	5 5-6 0		
Serum albumin	4 88		
Serum globulm (horse)	5.5		
Gelatin	4 80~4 83		
Castin	4 55		
Hemoglobin (reduced)	6 79-6 83		
Hemoglobin (oxidized)	0.7		
Ghadin	6.5		
Protammes	12 0-12,1		
Silk fibroin	2.0-2.1		
Myoain	6.2-6.6		
Pepsin	<1.0		
&Lactoglobulm	5 2		
Insulin	5 30-5.35		

EXPERIMENTS ON PROTEINS

- 1. General Composition Test. Heat some powdered egg albumln in a diftest tube in which is suspended a strip of moistened red litmus paper and across the mouth of which is placed a plece of filter paper moistened with lead acetate solution. As the powder is heated it chars, indicating the presence of carbon; the fumes of ammonia are evolved, turning the red litmus paper blue and indicating the presence of nitrogen and hydrogen, the lead acetate paper is blackened, indicating the presence of suffur, and the deposition of moisture on the side of the tube indicates the presence of hydrogen. (Moisture indicates hydrogen only in case both the powder and the test tube are absolutely dry.)
 - 2. Test for Organic Nitrogen—Lassaigne Test. Support a clean, dry test tube in a vertical position by means of a clamp placed near the open end. Drop in a small piece of freshly cut sodium (about a 3-mm. cube) which has been wiped free from oil with a piece of filter paper. Heat the bottom of the tube until a layer of sodium vapor about 1 cm. thick is formed. Drop a small amount of the material to he tested (enough to cover the end of a perknife or small spatula) directly on top of the hot sodium, avoiding contact with the upper wail of the tube. Heat strongly until most of the sodium has vaporized or until thick furmes cease to come off. When cool add a few milliters of distilled water. Heat the contents to boiling and filter, if the fusion with sodium has been carried out satisfactorily, the filtrate will be colories. Otherwise it should be rejected and the fusion repeated.

Boil 2 ml. of the clear, colorless filtrate for about a minute with a few drops of 19 per cent sodium hydroxide and 1 or 2 (no more) drops of a freshly prepared, saturated solution of ferrous sulfate. Cool, add dilute IICI, drop by drop, until the solution hecomes acid and the precipitate of ferrous hydroxide dissolves. The formation of a precipitate of prussian blue (ferra ferrocyanide) or of a hiue or green color in the solution indicates the presence of organic nitrogen in the original material. If the solution remains colorles add a drop or two of ferric chioride solution. If nitrogen is present the solution will turn hiue or green and a blue precipitate will usually form on standing.

The fusion of a nitrogen-containing organic compound with sodium results in the formation of sodium cyanide. This, when heated with ferrous sulfate in alkaline solution forms sodium ferrocyanide reacts with the ferric ion usually present, or with the added ferric ion, to form the blue ferric ferrocyanide. An excess of acid is to be avoided since the formation of the ferric ferrocyanide is much more delicate in the presence of only a slight access of acid to The reactions involved are as follows:

istic odor evolved from the solution. Write the reactions for this test. (2) Place equal volumes of KOH and egg-albumin solutions in a test tube and hoil the mixture vigorously. Cool, make acid with glacial acetic acid, and add 1 to 2 drops of lead acetate. A darkening indicates the presence of cysteine or cystine sulfur.

B. Test for Total Sulfill. (Cystine, Cystene, and Methionine). Place the substance to be examined (powdered egg albumin) in a small porcelain crucible, add a suitable amount of solid fusion mixture (sodium carbonate and potassium nitrate mixed in the proportion 2:1), and heat carefully until a coloriess mixture results. (Sodium peroxide may be used in place of this fusion mixture if desired.) Cool, dissolve the cake in a little warm water, and filter. Acidify the filtrate with hydrochloric acid, heat it to the boiling point, and add a small amount of harium chloride solution. A white precipitate forms if suifur is present. What is this precipitate?

As mentioned in the pieceding chapter, sulfur is piecent in proteins as cystine, cysteine, or methionine. Cystine and cysteine sulfur was formerly termed unoxidized, loosely combined, mercaptan, or lead-blackening sulfur. Methionine sulfur hecause of its greater stability toward alkaline lead acetate was called oxidized or acid sulfur. These terms are generally misleading and, in the case of the term oxidized, incorrect. The majority of proteins contain more methioniue S than cystine + cysteine S. Exceptions are keratins, insulin, and certain scrum albumins which contain all or almost all their sulfur in the form of cystine and cysteine. Silk fibroin and many protamiues are devoid of sulfur.

B. COLOR REACTIONS OF PROTEINS

These color reactions are due to a reaction between some one or more of the constituent radicals or groups of the complex protein molecule and the chemical reagent or reagents used in any given test. Not all proteins contain the same amino acids and for this reason the various color tests will yield reactions varying in intensity of color according to the nature and amount of the groups contained in the particular protein under examination. Various substances not proteins respond to certain of these color reactions, and it is therefore essential to submit the material under examination to several tests before concluding definitely regarding its nature.

1. Millon's Reaction. To 5 ml. of a dilute solution of egg albumin' in a test tube, add 3 to 4 drops of Millon's reagent.* Mix and hring the mixture gradually to the boiling point by heating over a small flame. Proteins like egg albumin, which are precipitated by strong mineral acids, yield a white precipitate which gradually turns red upon heating; whereas other proteins, like the secondary proteoses and peptones, yield only a red solution under the same conditions. If no color develops, add 2 to 3 more drops of the reagent

⁴ This egg albumin solution may be prepared by beating egg white with 6 to 10 volumes of water. The material is strained through cheescolds to remove the precipitate of non-mucin formed, and then filtered through filter paper and the filtrate used in the tests. A 1 per cent solution may also be prepared from poudered or scale egg albumin by soaking the material in a small quantity of water (sufficient to moster at throughly) for soveral hours, then diluting to volume, sterring until dissolved, and filtering
⁴ See Appendix.

and heat again An excess of the reagent is however in be avoided since it may produce a yellow color which is not a positive reaction. Repeat the test using a 0 i per cent solution of phenol instead of the protein, and note the red color produced upon iteating

This test is a particularly satisfactory one for use on solid proteins. In this case dilute some of the reagent with 3 to 4 volumes of distilled water add the solid and heat gently as above. The particles of indissolved protein will gradually turn red, if any of the protein dissolves the solution will also assume a red color.

The reaction is due to the presence of the hydroxy phenyl group —C.H.OH in the protein molecule and any phenolic compound which is unsubstituted in the 35 post tions such as tyrosine phenol (earholic send) and thymol will give the reaction Inasmuch as the tyrosine or haloscenated tyrosine grouping is the only hydroxy phenyl grouping which definitely has been proved to be present in the protein molecule it is evident that proteins respond to Million s reaction because of the presence of this amino acid. Tho test is not a very satisfactory one for use in solutions containing inorganic salts in large amount since the mercury of the Million's reagent is thus propertated and the reagent rendered inert. This reagent is therefore never used for the detection of protein material in the urine. If the solution under examination is strongly alkalinio it should be neutralized inasmuch as the alkali will precipitate yellow or black oxides of mercury.

2 Millan Nasse Reaction This is an adaptation of the Millon test and can be used in the presence of considerable quantities of inorganic salts, especially NaCi.

To 5 ml of a dilute solution mf pratein in a test tube, add 1 ml nf a 15 per cent solution of mercuric sulfate in 6 N sulfuric acid Place the tube in a boiling water bath for 10 minutes cool the contents in water for 5 to 10 minutes and add 1 ml of 1 per cent NaNO: A deep red color indicates tyrosine or other 35 unsubstituted phenol

3 Xanthoproteic Reaction To 2 to 3 ml of egg albumin solution in a test tube add 1 ml of concentrated nitric acid A white precipitate forms and upon heating turns yellow and finally dissolves, imparting to the solution a yellow color Cool the solution and carefully add ammonium hydroxide of sodium hydroxide in excess Note that the yellow color deepens into an orange Repeat the test using a 0 1 per cent phenol solution instead of the profuction, and note the production of the yellow and, later, the orange color

This reaction is due to the presence in the protein molecule of the phen) I group—Calls, with which the nitric and forms certain in tro modifications. The particular complexes of the protein molecule which are of especial importance in this connection are those of tyros ne and tryptophan. Phenylaha in does not respond to this test as it is ordinarily performed. The test is not a satisfactory one for use in urmary examination because of the color of the end react or.

4 Glyoxylic Acid Reaction (Hopkins Cole) Place 2 to 3 ml of egg albumin solution and an equal volume of glyoxylic acid (CHO COOH + H₂O or CH (OH), COOH) solution (Hopkins Cole reagent)* in a test tube and mix thor to flow slowly localize the tube and permit 5 to 5 ml of concentrated sulfure acid to flow slowly down the side of the tube thus forming a sharp layer of acid

See Appendix.

beneath the protein mixture. When stratified in this manner a reddishviolet color forms at the zone of contact of the two fluids. If the color does not appear after standing for a few minutes, the tube may be rocked gently to cause a slight mixing of the liquids at the interface. If the two liquids are mixed by gentle stirring the precipitate of protein dissolves and the violet color spreads throughout the solution.

This color is due to the presence of the tryptophon group (see p. 142). Gelatin does not respond to this test. Nitrates (NoNO, and KNOs), chlorates, nitrates, or excess of chlorides prevent the reaction, but a trace of copper sulfate will increase its sensitivity. The sulfuric acid used must be pure.

5. Birret Test. To 2 to 3 ml. of egg-albumin solution in a test tube add an equal volume of 10 per cent sodium hydroxide solution, mix thoroughly, and add a 0.5 per cent copper sulfate solution drop by drop, mixing between drops, until a purplish-violet or pinkish-violet color is produced. (If too much copper sulfate is added the violet color may be obscured by the blue precipitate of copper hydroxide formed.) The color depends upon the nature of the protein: proteoses and peptones give o decided pink; the color produced with gelatin is not far removed from a blue.

Repeat the birret test on some birret, formed from urea as follows: Place about one-eighth inch of urea in a clean dry test tube and heat gently over a small flame. The urea melts and then effervesces, and the birret formed appears as a white solid in tho bottom of the tube. Note the odor of the gas given off during the heating. Allow the tube to cool, dissoive the birret in 3 to 4 mi. of 10 per cent sodium hydroxide, ond add 0.5 per cent copper suifate solution, drop by drop, until the pink color oppeors.

The bluret test may also be carried out with o stablo bluret reagent, prepared by adding 1 per cent copper sulfate solution, drop by drop, with constant stirring, to some 40 per cent sodium hydroxide solution until the mixture assumes a deep blue color. This reagent may be used in either of two ways. It may be added directly to the protein solution, o drop at a time, with mixing, until the solution assumes o violet color; or two or three drops of the reagent may be permitted to flow down the side of the inclined tube. In this case the reagent forms a layer henceth the protein solution and the violet color appears at the interface between the two ilquids.

The bluret test is given by those substances whose molecules contain two earbampl (—CONII-) groups joined either directly together or through a single atom of introgen or earbon. Similar substances which contain (so place of the —CONII-, group) —CSNII-, —C(NII)NII-, or —CH₂NII-, also respond to the test. It follows from this fact that substances which are nonprotein in character but which contain the necessary groups will respond to the hieret test. As examples of such substances the following may be etted:

CONII. CONH.
CONII. NII
CONII.

According to Schiff the end reaction of the burset test is dependent upon the formation of a copper potassium burset compound (eupripotassium burset or burset potassium eupric hydroxide). This substance was obtained by Schiff in the form of long red needles and has been formulated as follows.

Haurowitz' assumes that a copper coordination complex with the following ring structure is probably produced

If much magnesium sulfate is present a precipitate of magnesium hydroxide forms which interferes with the test. If much ammonium sulfate is present a large excess of alkali must be used.

6 The Triketohydrindene Hydrate (hinhydrin) Reaction To 5 ml of dilute protein solution, which must be approximately between pll 5 and pll 7, add 0.5 ml of a 0.1 per cent solution of triketohydrindene hydrate heat to boiling for one to two minutes, and allow to cool A blue color de velops if the test is positive

This test gives positive results with proteins, peptones, peptides amino acids, and other primary amines, including aminonia For further discussion see p 129

C PRECIPITATION REACTIONS OF PROTEINS

The proteins are precipitated from solution by salts of the beaty metals (e.g. HgCl: \g^0\), CuSO, Pb(C4H₂O₂), etc.), by certain acids some of which are called all abudal reagents (pience acid, phosphotungstic acid tannic acid metaphosphoric acid etc.) by concentrated solutions of such salts as ammonium sulfate sodium sulfate and sodium chloride and by ethyl and methyl alcobol. Although these reactions have been used for many years for the separation and characterization of proteins there is still no definite evidence concerning the nature of the mechanisms.

¹ Haurowitz Chemistry and Biology of Proteins New York Academic Press, Inc. 1950 p. 11

t A few drops of pyridins or a few crystals of sodium acetate may be used to adjut the

involved Some of this uncertainty is due to the experimental difficulties involved in the isolation of the pure products formed in these reactions. There is also a great deal of confusion due to attempts by various authors to interpret these reactions exclusively on either a chemical or a colloidal basis.

In the case of the acids, the evidence suggests that the protein comhines with the acid radical to form insoluble salts, such as protein tannate, protein phosphotungstate, etc. This is in agreement with the views of Loch, who pointed out that proteins combine with negative, or acid, ions only on the acid side of their isoelectric points. In agreement also is the fact that if, instead of the free acids, the salts of these acids are added to neutral solutions of protein, no precipitates result until the solution is acidified Some of the reactions obtained with the salts of the heavy metals are probably similar in character, precipitates of silver proteinate, for example are formed on the alkaline side of the isoelectric points. The reactions are, however, complicated by the fact that in some cases the first addition of salt causes the formation of a precipitate which dissolves in excess of salt and then reprecipitates when more salt is added. The difficultics involved in arriving at a rational explanation of these reactions are intensified by the fact that individual proteins show great differences in hehavior, probably because of differences in the hydration and dispersion of the particles in solution, denaturation, etc. In general, precipitants of this class usually cause extensive intramolecular changes in the protein molecule

The proteins, in common with other emulsoids, are precipitated by concentrated solutions of such salts as ammonium sulfate, sodium sulfate, sodium chloride, etc. In these cases, as discussed in Chapter 1, precipitation is apparently due to the neutralization and dehydration of the molecules and molecular aggregates in solution. The protein precipitated by these silting out methods is unaltered (native) and usually redissolves when treated with fresh portions of the original solvent. The concentration of salt required for the precipitation of a protein depends on the particular protein and on the pH of the solution—i.e., on the charge on the protein complex. These variations will be considered in the next chapter in connection with the properties of the various classes of proteins.

The proteins are also precipitated by dehydrating agents, such as alcohols and acctone. The addition of alcohol to electrolyte-free solutions of proteins converts them into suspensoids, which floculate upon the addition of a few drops of salt solution. Precipitation by alcohol is most effective at the isoclectric point of the protein. Besides precipitating proteins, ilcohol acts on certain proteins such as egg albumin, to produce intrinsolucidar changes which affect the solubility and other properties of the protein. This phenomenon, known as denaturation, will be discussed more fully in the next section. The denaturing effect of alcohol on some proteins may be minimized by the use of temperatures around 0° C or below. Prolonged contact with alcohol (or any other denaturing agent, for that matter) produces an increasible coagulation of the pre-tiem. The fixing of tissues for histological examination is an example of the coagulating action of alcohol on proteins.

Colloidal iron, kaolin, and alumina cream are frequently used for removing proteins from solution. These substances probably act by advorption and their use has been adapted to various quantitative methods.

1 Effect of Strong Acid and Alkalt Place a few ml of concentrated nitric acid in a test tube, incline the tube, and add dilute egg albumin slowly from a pipet, allowing the solution to run down the side of the tube and form a layer over the nitric acid. Note the appearance of a protein precipitate at the zone of contact between the two fluids. Now mix the contents of the tube thoroughly by careful shaking is protein precipitated by concentrated nitric acid?

Repeat the above experiment using concentrated sulfuric acid, concentrated hydrochloric acid, acetic acid, and concentrated sodium hydroxide. How do these various reagents differ in their action on proteins? Allow the tubes to stand overnight or longer and note any further changes.

The formation of a protein precipitate by layering the solution over nitricacid as described above in frequently used as a test for protein in urine and other fluids (Heller's test, see p 830)

2 Precipitation by Metallic Saits Prepare six tubes each containing 2 to: ml of dilute egg alhumin solution To the first add mercuric chloride solution, drop by drop slowly, until an excess of the reagent has been added, as noteany changes which may occur Unless the reagent is added very gradual the formation of the precipitate may not he noted, due in its soluhlity access of the reagent Repeat the experiment with lead accetate, allver nitraticopper sulfate, ferric chloride, and barium chloride, using very dilute solutions.

Egg albumin is used as an antidote for lead or mercury poisoning Why? Is it an equally good antidote for the other metallic salts tested?

- 3 Precipitation by Alkaloidal Reagents Prepare six tubes, each cootaining 2 to 3 ml of dilute edg albumin solution. To the first add pictre acid, drop hy drop, until an excess of the reagent has been added, noting any changes which may occur. Repeat the experiment with trichloroacetic acid, tannic acid, phosphotungstic acid, phosphotungstic acid, phosphotungstic acid, and potassio mercuric lodide. Are these precipitates soluble in excess of the reagent? Acidify with hydrochloric acid before testing with the last three reagents.
- 4 Precipitation by Ferrocyanide To 5 ml of dilute egg albumin solution in a test tube add 5 to 10 drops of acetic acid. Mix well and add potassium ferrocyanide, drop by drop, until a precipitate forms. This reaction is very sensitive.

solution is saturated with the salt. What happens? Again dilute a portion of the mixture with water. Is this precipitation reversible? Fifter the remainder of the mixture and test a portion of the precipitate by the Millon test. Test the filtrate by the bluret test, using a saturated solution of ammonium sulfate as a control and adding the same amounts of alkali and copper sulfate solution to hoth control and filtrate. What are your conclusions?

(b) Repeat the above experiment, using sodium chioride instead of ammonium sulfate. How do the results differ from those obtained with ammonium sulfate? At the saturation point with sodium chloride, add 2 to 3 drops of acetic acid. What occurs?

All proteins except peptones are precipitated by saturating their solutions with ammonium sulfate Most globulins are precipitated by halfsaturation with ammonium sulfate or full saturation with sodium chloride. If the saturated sodium chloride solution is subsequently acidified, all proteins except peptones are precipitated

Soaps may be salted out in a similar manner (see p 108).

6. Precipitation by Alcohol. (a) INFLUENCE OF ELECTROLYTES Prepare an electrolyte-free solution of egg albumin as foilows. Place the albumin solution in a dialyzing bag (see p. 10), add a drop of toluene as apreservative, and tie the mouth of the bag securely. Immerse in a large beaker of distilled water and allow to stand for several days, changing the water at intervals. Remove the albumin solution from the bag and filter. Test the filtrate for chloride, it should be negative; if not, the dialysis must be repeated.

Place a 5-mi. portion of the salt-free albumin solution in each of two test tubes and add 10 ml. of 95 per cent alcohol to each tube. Mix. Now to one tube add a pinch or two of solid sodium chloride and again mix. Compare results in the two tubes. What is the effect of alcohol on protein in the absence of electrolytes? What is the effect of added electrolyte? Explain. Dilute some of the suspension with water. Does the precipitated protein redissolve?

(b) INFLUENCE OF ISOELECTRIC POINT See Exp. 2, helow.

D. ISOELECTRIC POINTS OF PROTEINS

1. Isoelectric Point and Solubility of Casein. Into a 50-ml. volumetric flash Introduce 0.25 g. of pure casein. Add about 20 ml. of water and exactly 5 ml. of N NaOH. When solution is complete add exactly 5 ml. of N acetic acid and dilute to 50 ml. Vix well. This is a solution of casein in 0.1 N sodium acetate. Set up a series of nine tubes as follows.

Tube No	1	2	3	4	5	6	7	s	9
Distilled water mil 0 01 N acetic acid mil	S 38 0 62	7.75 1 25	8 75	8 5	8	7	5	1	7 4
1 0 N nectic held mil			0 25	0.5	1	2	4	8	1 6

indicating no turbidity by θ , degrees of turbidity by + signs, and degrees of precipitation by \times .

Tube No	1	2	3	4	5	6	7	8	9
pli Turbidity, immediate Turbidity after 5 minutes	5 9 0 0	5 6 0 0	5 3 + +	++	4 7 +++ ×××	4 4 ++ ××	+	3 8 + +	3 o 0 0

The precipitation should be greatest in Tube 5 which has a pill of 47, near the isoelectric point and point of least solubility of casein. The aclist in each tube may be actually determined by the electrometric method or may be calculated from the concentrations of sodium acetate and acetic acid by means of the Henderson-Ilasselbaich equation (see p. 33).

2 Isoelectric Point and Precipitation of Gelatin by Alcohol. Gelatin and many other proteins are quite soluble in water even at their isoelectric points They do, however, precipitate more readily at this point if some precipitating agent such as alcohol is added Prepare a series of test tubes as follows

Tube Vo	1	2	3	4	5	б	7	8	9
01 N sodium acetate ml 01 N acetic acid ml 10 N acetic acid ml 10 N acetic acid ml Distilled water ml 17, gelatin ml	2 00 0 12 3 88 2 00	2 00 0 25 3 75 2 00	100	•	1 2	4		16	3

Mix the contents of the tubes well and add 95 per cent alcohol to Tube 5 until a very faint cloudiness is produced (ordinarily about 8 ml are required). Add the same amount of alcohol to each of the other tubes Examine after 30 mlnutes. The results usually obtained, with approximate pil for each tube, are as follows:

	-							
Tube \a	i 1	2	3	4	5	G	~	8 9
						-	_	
Turi stity			-	++	+++	+	- ·	- 1 -
pH	6.0	56	5 1	5 0	1.7	4.4	3 1	38135

The Isoelectric point for gelatin is about pil 47

of great blological importance Relatively slight changes in pH greatly influence protein swelling

Prepare a series of test tubes as follows

Tube \o	1	,	3	4	J	в	7	8	9	10
10 \ HCl ml 01 \ HCl ml Distilled water ml	16	8 12	16	20 0	10 10	5 15	2 o 17 o	1 2a 18 7a	0 62 19 38	0 31 19 6)

Gut strips as nearly the same size as possible ($5\,\mathrm{cm} \times 0.5\,\mathrm{cm}$) from a sheet of gehatin at least 1 mm thick. Fut a strip in each tube. After 24 hours measure the length of the strips. Results similar to the following may be obtained

Tube No	1	.3	3	4	0	6	~	s	9	10
I ength (cm)	6 4	6 6	67	7 4	83	83	7 6	7 0	6.3	5 3

The greatest degree of swelling will usually be found in tubes containing from 0.025 to 0.05 N HCl corresponding to pH salues of 1.3 to 1.6 The reaction within the gelatin particles themselves at the point of maximum swelling is, however, about pH 3.2 This difference in reaction within and without the gel is a consequence of the Donnan equilibrium (see p. 12) which must be considered in all such cases involving the relations of protein ions to other ions in solution. Instead of gelatin strips, 1 g. portions of powdered gelatin (between 30 and 50 mesh) may be placed in 100 ml. graduated cylinders containing suitable solutions and the relative degree of swelling measured. The actual pH of the gel in any case may be determined electrometrically.

A Influence of Acidity on Solidification of Gelatin Tho isoelectric point for gelatin (pH 47) is the point of minimum viscosity of its solutions and the point at which the gelatin most readily solidifies This is because at this point the gelatin is least soluble in the more liquid (water) phase an 1 hence imparts itse viscosity to it For the same reason there is a greater tendency for the more solid phase (gelatin) to form the supporting network believed to be the basis of a jelly Viscosity determinations must of course be made on the liquefied gelatin solution

Prepare a series of test tubes as follows

Lube Vo	t	,	3	4	٥	G	-
01 \ so hum rectate mt 01 \ sectic icid mt	1 00	1 00 0 06	1 00 0 25	10	10	1 0	1 0
10 N netic cilni 10 N so hum hydravide i i Water i i	0 0s	 6 91	b 7a	60	3 0	16	06
pli (apį n v)	9 00	5 (0	a 00	46	4 0	3 4	2 8

To each tube add 3 ml of 4 w rm 10 per cent solution of gelatin and mix place ill the tubes in water at 50°C for a few minutes, then transfer to a large beaker of water at room temperature, noting the time at this point After a few minutes in the beaker, the tubes may be removed and placed in a rack to facilitate observation. Note the time required for solidification of the gelain in each tube. Degree of solidification may be established by thiting the tube and observing whether or not the fluid will flow. Results similar to the follow ine should be obtained.

with a second se							
Tube No	1	2	3	4	5	6	7
Time required for solidification (mm)	19	17	16	14	19	25	long
	1		ļ	·			

Observations may be repeated by remelting the jellies Note also the much greater opacity of the gelatin at the isoelectric point, indicating that most of the gelatin is in the solid phase

E PROTEIN DENATURATION AND ITS REVERSAL

A protein is called a native protein if its amino acid composition and stereochemical structure are unchanged from the natural state. These properties control all the functions of a protein, whether solubility is dilute sait solutions, proteolytic activity, oxygen-carrying capacity, or whatever it may be. These characteristics are altered and the process of denaturation is said to occur when a protein undergoes changes in structure or composition. Chemical and physical agents which cause these changes are called denaturing agents. Their action involves the splitting of some or all of the protein cross linkages with their possible re-formation in some cases, to cause a rearrangement of the peptide chains.

It is to be expected that proteins will vary widely in their ease of denaturation Certain proteins especially those which in solution are fibrous ur highly clongated molecules such as the musclo protein myosin, are easily denatured, others, such as the earbohydrate-rich glycoproteins (oxomucoid, for example) seem to be quite resistant to the usual physical

arents causing denaturation

Denaturation may be caused by (1) heat which causes splitting of the salt bridges by thermal agitation (2) mineral acids and alkalies, which alter the ionization of carboxyl and amino groups respectively and thus destroy salt bridges (3) shaking or stirring which results in the dena turation of the protein spread over the surface of the air bubbles (4) grinding, which probably causes mechanical deformation of the peptide chains (5) ultraviolet radiation which splits the peptide bonds adjacent to the arom use rings (6) ultrasonic waves which destroy the rings of the aromatic amino acids and whose denaturing power depends on their friquidity (7) incutral chemical agents such as urea and certain acid and guandline dividuous that act as dispoles to cause (Lavage of hydroga bonds and formation of new ones * It has been subjected "that the brief and of hydrogan bonds hy urea chables the "all groups to react with intramolecular". So bridges stock of which the second of the mean of the stock of which the second of the subjected of the property of the intramolecular.

an intermolecular link with the liberation of a new —SH group. The coutinuation of this chain process gives rise to an —S—S— cross linked framework of higher molecular weight and decreased solubility at the isoelectric point. This reaction is formulated as follows.

Denaturation then, appears to be an all or none phenomenon that is, a protein is either denatured or unchanged from its natural state. Denatured proteins may, under certain conditions be restored to proteins ath many of the properties of the original protein. This is known as cenaturation. It appears however that renaturation seldom results in the complete transformation of the denatured protein to its original state. Thus we may classify proteins into three groups natural denatured and renatured.

If the foregoing rather specific definition of denaturation is used then as already stated protein deusturation must have an all or none character However, denaturation seldom is completed in one step but consists of a series of chemical and structural deviations from the original native protein If these changes are not too extensive many of the prop erties possessed by the native protein may be restored. Thus pepsin can be denatured and so lose its protectivity properties if it is warmed to the proper temperature When the solution is cooled the proteclytic activity of the protein is restored-i e denaturation is reversed and renaturation results Likewise, the oxygen enriging capacity of hemoglobin can be destroyed by denaturing with salieylate On reversal the restored hemoglobin is very similar to the original untreated hemoglobin. Although many of the properties of the native protein can be restored by reversal regen eration, it is still question ible in many instances whether the reversed protein is identical with the original native protein. Many regenerated proteins may simulate certain characteristic properties of the native materials from which they were derived but still remain denatured proteins

It is thus apparent that the extent of denaturation and its reversal is primarily dependent upon the methods used Probably the oldest means of denaturing a protein and one which is familiar to all is illustrated by the marked change in consistincy of egg white on cooking, a process which results in extensive denaturation of the albumen proteins. What is seen when egg white is cooked is only the end result of denaturation—the matting together of the fibers of denatured protein to form a tightly adhering coagulum. Den iteration involving less deep-scated changes in protein structure requires more refined methods to detect. The studies of

Roche, Neurath, and others have shown that denatured proteins which other investigators believed to have been entirely reversed to the native form were still denatured when more delicate methods of measurement were employed

In practice, then denaturation consists of a series of changes in the protein molecule brought about by various chemical and physical agents. These changes often affect the viscosity, particle size, solubility, restance to proteoly tic enzymes and formation or disappearance of sulfhydrigoups, and may even cause the loss of certain amino acids or peptides of relatively low molecular weight.

Denatured proteins, because of their reduced solubility, usually fleculate at or near the isoelectric point. This flocculation is ordinarily reversible at room temperature. However, if the suspension at the isoelectric point is heated, the floccules form relatively large, tenacion mass, or of coagulated protein which are not easily redused by treatment with dilute acids or alkalies. Denaturation is the primary and important change, flocculation and coagulation, which were often confused with dilute acids or alkalies. Denaturation of the changes in the protein molecule brought about by denaturation.

1 Denaturation Flocculotion and Coagulation of Egg Albunin Place 9 ml of a clear, salt free solution of egg albumin into each of three test tubes 70 the first add 1 ml of 01 N liCl, to the second 1 ml of a mixture of solum acetate and acetic acid (pil 47) and to the third 1 ml of 01 N \aOH llest all three tubes in a bolling water bath for 15 minutes Cool and examine 70 tubes 1 and 3 add 10 ml of acetate buffer solution, pH 47 What happens litter off the precipitates in each tube and wash them on the filter papers with distilled water Precipitates 1 and 3 are denatured egg albumin Precipitate 2 is coagulated egg albumin Precipitate 2 is coagulated egg albumin

Suspend each of the precipitates in about 10 ml of distilled water and dilvide each suspension into three parts. To the first part add dilute HCl, drop by drop. Does the precipitate dissolve? Repeat with another part of the suspension, adding dilute \aOll Heat the third parts of suspensions 1 and 3 ln a bolling water bath for 15 minutes. Cool and test the solubility of the material in dilute acid and alkall Does the material now dissolve?

The precipitates of denatured protein formed by bringing their solutions to the isoelectric point dissolve readily in a few drops of dilute acd or alkali. The coagulated protein remains insoluble under similar conditions. Heating the precipitate of denatured protein converts it into a coagulated protein in which case it exhibits the properties characteristic of that type of substance.

F IMMUNOROGICAL REACTIONS OF PROTEINS

Certain materials when injected into a bying animal under proper conditions stimulate the animal to produce substances which have the power to react with the specific material injected. The substance injected is referred to as the anti-pin while the specific substances appearing in the blood of the immunized minimal are known as antibodies. In order to act as anti-constitue the substances must be protein for combinable with protein for combinable with protein for combinable with protein for combinable with protein for combinable with protein for combinable with protein for combinable with protein for mature collected and foruga to the blood stream of the animal injected. The antibodies are structured.

ically) modified serum proteins, usually globulins, induced by the presence of the antigen in those organs in which the formation of serum proteins occurs. Their presence in the blood stream of the immunized animal is recognized by their action on antigens. The type of reaction which takes place between the antigen and antihody depends upon the nature of the antigen. If, for example, the antigen consists of a suspension of red blood corpuscles, the blood serum of the immunized animal, known as the antiserum, acquires the property of hemolyzing this particular type of corpuscle. If the antigen is a suspension of hacteria or other cells, the antiserum will cause these particular cells to clamp together or agriptionals.

If the antigen is a soluble protein the antiserum is capable of forming a precipitate with this specific protein, when mixed under proper conditions. This reaction, known as the precipitin reaction, is remarkably delieate and specific By means of the precipitin reaction it is possible, for example, to detect a specific protein in solutions containing as little as one part in ten million, or even in one bundred million Furthermore, this reaction is so specific that the antiserum prepared by injecting a solution of heu's egg albumin will precipitate ben's egg albumin but will give no reactions at extreme dilutions with egg albumins from other sources, or with any of the other proteins in egg white. The precipitin reaction is thus of great value in determining the identity and individuality of protein preparations. It permits us to detect differences between similar proteins from elosely related sources which, in their elemical composition and properties, are practically indistinguishable. By means of this reaction it may be shown that certain proteins, such as the protein of crystalline lens are immunologically the same in animals of widely differing species. It has been used to study relationships between various species of plants and of animals Practically, it is used in medicolegal work to distinguish specifically between human and other bloodstams

The technique involved in immunization and in carrying out precipiting reactions is illustrated in an experiment on p. 485

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Proteins: Their Classification and Properties

The classification of the proteins is a very difficult problem because, as was outlined in Chapter 5 they are found in nature not as distinct chemical individuals but as parts of larger complexes formed by the union of various proteins with each other and with carbohydrates and lipides. The problem is further complicated by the fact that individual purified proterns may themselves be separated into components showing individual differences in composition and properties. Until these components are isolated in pure form and their exact structure determined, a classification of proteins based on molecular structure is impossible. The classification offered below is based upon the fact that under specified experimental conditions the proteins in biological materials (e.g. egg white or blood pla ms) msy la separated into groups which exhibit certain characteristic properties It should be kept in mind however, that these proteins probably do not exist as such in the native material, and that, furthermore their composition and properties are dependent upon the methods used in their relation. In spite of all its shortcomings such a classification permits the grouping tegether of proteins with similar properties into fairly definite classes and must suffice until the information required for a more rigid clas incution is obtained

CLASSIFICATION

I SIMPLE PROTEINS

Albumins are soluble in salt-free water and coagulable by heat, e.g., ovalbumin from egg white, scrum albumin from blood serum, lactalbumins from milk. vegetable albumins

It is now recognized that serum albumin is not a homogeneous material but can be fractionated by salting-out procedures (cf. pp. 163-164) into a number of distinct components all of which have albumin characteristics. The term serum albumins is preferable to the more widely used serum albumin. These proteins are, however, either not separated or but poorly separated by the widely used ultracentrifuge or electrophoretic methods (See p. 461).

Globulins are insoluble in salt-free water but soluble in neutral solutions of salts of strong bases with strong acids, such as NaCl, and are coagulable by beat, e.g., serum globulin, lactoglobulin, thyroglobulin, edestin from hemp seed, amandin from almond, and other vegetable globulins

Serum globuln has been fractionated into several components by ultracentifuge and electrophoretic methods (see Chapter 22). The three groups most castly identified are called α , β , and γ globulins. The γ -globulin fraction of buman plasma is the source of almost all the immunologically active proteins of the blood. In fact, it appears that 50 per cent or more of the γ -globulin fraction prepared from pooled human plasma is composed of immunologically active material. During convalesceuce from illness, there is an increase in immune proteins in the plasma. This increase is accompanied by and accounted for by the increase in γ -globulin. Figure 57 shows a paper electrophoretic pattern of normal buman serum

Glutelins are simple proteins insoluble in all neutral solvents, but readily soluble in very dilute acids and alkalies, e.g., glutenin from wheat

These may be mixtures of denatured proteins

Prolamins are simple proteins soluble in 70 to 80 per cent alcobol and insoluble in water, absolute alcohol, and other neutral solvents, e.g., zen from corn, gladin from wheat and ryc, bordein from barley, and hynin from malt. Rice, hafir, and sorgbum bave also been shown to contain alcohol-soluble proteins. The name prolamins was suggested for these alcobol-soluble vegetable proteins by Osborne, since upon by drolysis they yield large amounts of proline and ammonia.

Albuminoids (Scleroproteins) comprise all fibrous proteins which have a supporting or protective function in the animal organism. In the plant langdom the albuminoids are probably replaced by cellulose and similar substances.

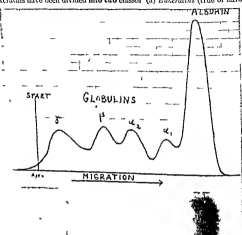
I COLLAGE'S, the principal supporting proteins of skin, tendons, and bones, are resistant to peptic and tryptic digestion. They are converted into the easily digested soluble proteins, the gelatins, by boiling with water, dilute aeids, or alkalies and contain large amounts of hydroxyproline.

2 ELASTINS, present in elastic tissues (tendons, arteries), are more readily digested by trypsin than by pepsin, are not convertable into gelatin, and give a negative or very weak test for hydroxyproline Clustic tissue is a mixture of elastin, collagen, and a carbohydrate-containing protein, clastomicin It is probable that many of the earlier studies

on "clistin" were really investigations on a mixture of clastin and clastomurin ²

3. At RATINS are proteins resistant to digestion by pepsin and trypsin

3 Kerkkins are proteins resistant to digestion by pepsin and trypsin and involuble in dilute acids and alkalies, in water, and in organic solvents Keratins have been divided into two classes (a) Euleratins (true or hard



yield a large number of amino acids, among which arginine predominates Histones contain tyrosine hit appear to be lacking in tryptophan In short, bistones are basic proteins which stand between proteinines and true proteins, e.g., scombrone from mackerel sperm, thymus histone

Globins are simple hasic proteins which were at one time classed as histones because of similarities in solubility, etc. The globins differ from the bistones, however, in isoelectric point, toxicity, and amino acid composition. Histones are high in arginine and isolectine and contain only traces of tryptophan, but globins contain average quantities of arginine and tryptophan and are unique in their high content of histidine and deficiency in isoleucine. Globius are usually found in nature as the protein portion of conjugated proteins, e.g., globin from hemoglobin.

Protamines are simpler polypeptides than the proteins included in the preceding groups. Their acid salts are soluble in water, uncoagulable by heat, bave the property of precipitating aqueous solutions of other proteins, possess strong basic properties, and form stable salts with strong mineral acids. They yield comparatively few amino acids, among which the hasic ones, especially arginine, predominate. They are the simplest natural proteins e.g., salmine from salmon sperm sturine from sturgeon sperm, clupeuie from herring sperm scombrine from mackerel sperm.

II CONJUGATED PROTEINS

The conjugated proteins, like the simple proteins, are substances which may be isolated from biological materials by suitable methods. They differ from the simple proteins however, in the fact that the intact molecule consists of protein combined with some nonprotein substance (the prosthetic group) in a manner which confers new and characteristic properties out the complex formed. They are classified according to the nature of the prosthetic group, as indicated below.

Nucleoproteins. Compounds of one or more protein molecules with nucleic acids. The nucleic acid of the cell nucleic suppears to be decrynthese nucleic acid which is united to the protamnes histones, and other base proteins of the cell nucleis. The nucleoprotems of the cytoplasm yield abose nucleic acid, the nature of the protein component awaits investigation.

Glycoproteins. Mucins contuin a uronic acid (see Chapter 2) probably united in salt liukage to the basic groups of proteins. They are found in vitreous humor and in Wharton's jelly of the umbilical cord

Mucoids do not contain a uronic acid but consist of protein firmly bound to a polysacchride such as polymenzed glucosamine-mannose, e.g., serum mucoid, ovomucoid

Sulformacins contain sulfuric read, urome acid, and either chondrosamine or glucos imme. They are found in earthlage, intestinal tissue, cornea, gastric mucosa.

Phosphoproteins. Compounds of the protein molecule with phosphoric acid other than a nucleu acid or lecithur, e.g., casein from milk, ovoritellin from egg yolk, and other proteins associated with the feeding

The accumulated chemical evidence distinctly points to the propriety of classifying the plosproteins as conjugated compounds i.e. they are esters of phosphoric acid (or acid) and rotein

of the young In the cell nucleus there are phosphoproteins the composition of which is still under investigation

Chromoproteins are compounds of proteins with a metal-containing prosthetic group, e.g. the red iron containing hemoglobins (Fe = 0.34 per cent) from vertchrate blood the blue copper containing hemocyanins from the blood of certain invertebrates, the green magnesium-containing chlorophyll proteins from plants etc. The protein should not be regarded as a mere colloidal carrier of the prosthetic group, but as an integral portion of the chromoprotein molecule determining not only the magnitude but even the nature of the reaction it promotes. Thus one iron-containing prosthetic group combined with four different proteins gives rise to four different substances methemoglobin, eatalase, peroxidase, and a cytochrome ($\Gamma e = 0.43$ per cent)

Lipoproteins are compounds of the protein molecule with lecithins,

Metalloproteins are various protein enzymes which contain metals as an inherent portion of their molecule Thus, tyrosinase contains copper, arginaso contains maoganese or magnesium whereas zine is present in carbonic anhydrase, and molybdenum in xanthine oxidase

III DERIVED PROTEINS

PRIMARY PROTEIN DERIVATIVES

The primary protein derivatives are substances formed from some of the simple and conjugated proteins on denaturation. When these proteins are subjected to certain chemical and physical agents they undergo intramolecular changes which are accompanied by changes in the properties of the original material Although the nature of these intramolecular changes is still unknown, they may be recognized by the characteristic properties of the substances formed as indicated in the classification

Proteans are insoluble products which apparently result from the inespicint action of water, very dilute acids or enzymes-eg, fibrin from fibrinogen, myosan from myosin cdestan from edestin. This is probably an carly denaturation stage

Metaproteins are products of the action of dilute acids and alkalies whereby the molecule is so far altered as to form products soluble in weak acids and alkalies but insoluble at their isoelectric points e.g., acid metaprotem (acid albuminate) alkalı metaprotein (alkalı albuminate)

Coagulated proteins are insoluble products which result when isoelectric solutions of the protein are denatured by (1) heat, (2) alcohol, (3) ultraviolet light (4) ultrasonie waves (5) mechanical shaking (6) mineral acids and alkalics, (7) grinding (8) neutral chemical agents, etc

SECONDARY PROTEIN DERIVATIVES

The secondary protein derivatives are substances formed during the hydrolysis of the protein molecule As hydrolysis proceeds the intact

Tile term secondary protein derivatives is used because their formation may be preceded by that of some of the primary derivatives.

molecule is split up into a series of smaller and smaller fragments which are designated respectively, as primary and secondary protoces peptiones, and peptides. The substances in each group are not homogeneous chemical entities, but rather mixtures of fragments of the original protein which probably vary in both composition and size and which are grouped together metely because they exhibit certain characteristic properties in common. This classification has little modern significance and it is being gradually abandoned except for the peptides.

Primary proteoses are soluble in water, noncoagulable by heat, precipitated by concentrated nitric acid, and precipitated by half-saturation

with ammonium sulfate

Secondary proteoses are soluble in water, noncoagulable by heat, and precipitated by saturating their solutions with ammonium sulfate

Peptones are soluble in water noncoagulable by heat and not precipi-

tated by saturating their solutions with ammonium sulfate

Peptides are definitely characterized combinations of two or more amino acids, the carboxyl group of one being united with the amino group of the other with the elimination of a molecule of water, 6—e g, dipeptides tripeptides, tetrapeptides, pentapeptides, etc.

CONSIDERATION OF VARIOUS CLASSES OF PROTEINS

I SIMPLE PROTEINS

A ALBUMINS

Albumms constitute the first class of simple proteins and may be defined as simple proteins which are coagulable by heat and soluble in pure (salt-free) water. They are also soluble in salt solutions and those of animal origin are not precepitated upon saturating their solutions at 30°C with sodium ebloride or magnesium sulfate except in the vieinity of their isoelectric points. All albumins of animal origin are soluble in half-saturated ammonium sulfate solution? but may be precipitated by increasing the salt concentration up to full saturation. They may also be thrown out of solution by the addition of a sufficient quantity of a mineral acid Metallic salts also possess the property of precipitating albumins, some of the precipitates being soluble in excess of the reagent, whereas others are insoluble in such an excess. Many albumins have been obtained in crystalline form, notably egg and serum albumins from various species.

EXPERIMENTS ON ALBUMINS

Besides the general protein color reactions and precipitation reactions described in the previous chapter the albumins have other properties which are used to identify proteins belonging to this class. Some of these properties are illustrated by the following experiments

⁶ The peptones are undonbtedly peptides or mixtures of peptides the latter term being at present used to de ignate those of definite structure

In this connection Osborne's observation is of interest namely that certain vegetable albumins are pre-instact by saturating their solutions with sodium chloride or magnesium sulfate or by half saturating with ammonium sulfate

- I Solubility in Concentrated Salt Solutions (a) Place 25 ml of dilute (££ albumin solution (prepared as described on p 169) in a beaker and add 6 6 ½ of solid ammonium sulfate. Stru until dissolved. The solution is now 2 M in ammonium sulfate, or approximately half saturated. Is the albumin precipitated? Now add an excess of solid ammonium sulfate and still to produce full saturation with the salt. What happens? Didute a small portion of the mixture with distilled water. Is the effect of ammonium sulfate reversible? Filter the remainder of the mixture. Test the precipitate by Millon's test. Test the filtrate for protein by the bluret test, using saturated ammonium sulfate solution as a control What are your conclusions?
- (b) To a 25 ml portion of eag albumin solution in a beaker add an excess of solid sodium chloride and stir until the solution is saturated with the sait How does the result differ from that with ammonium sulfate? Add 2 to 3 drops of acetic acid. What occurs? Explain
 - 2 Heat Coagulation Add 3 to 5 drops of acetic acid to 25 ml of dilute egg albumin solution in a small evaporating dish and heat to boiling. What happens? Why is the acetic acid added? Test a portion of the coagulum by Millon's reaction. Explain

B GLOBULINS

Globulins are simple proteins present in blood serum, muscle, and other asimal tissues and also present in many plant seeds. They give all the ordinary protein tests and are coagulable by heat Globulins differ from the albumins in being insoluble in pure (salt-free) water. They are, however soluble in neutral solutions of the salts of strong acids with strong bases such as sodium chloride. The globulins require a certain concentration of salt in order that they may remain in solution precipitating when the concentration of salt is lowered by such processes as dilution or dialysis. In general the globulins are precipitated by half-saturation of their solutions with ammonium sulfate—i.e. by the addition to their solutions of an equal volume of saturated ammonium sulfate solution. Most globulins are also precipitated from their solutions by saturation with solid sodium chloride or magnesium sulfate.

Blood serum apparently contains a variety of globulins, characterized by differences in solubility in precipitability by ammonium or sodium sulfate, and in rate of cleetrophoretic migration (Chapter 22) Blood globulins have not yet been crystallized but crystalline globulins have been obtained from milk (6 lactoglobulin) muscle (myosin), gastric juice (pessin), and numerous plant, seedle

It is generally stated that globulus are precipitated from their solutions upon I all saturation with amongum sulfate and that albumins are precipitated upon complete saturation by this said. Comparatively and that albumins are precipitated upon complete saturation by this said. Comparatively and the said of vegetable original cases to be more extensively dided These studies furthered especially by Oblome and associates have demonstrated all all resultants of the three controls of the said of

We have used an alhumin of animal origin (egg albumin) for all the protein tests thus far, whereas the globulin to be studied will he prepared from a vegetable source The vegetable globulin we shall study may be taken as a type of all globulins, both animal and vegetable.

EXPERIMENTS ON GLOBULIN

Preparation of Edestin. Extract 20 to 30 g. (a handful) of crushed hemp seed with about 150 ml. of 10 per cent sodium chloride solution for one-half hour at 60° C. Filter while hot through a filter paper moistened with 10 per cent sodium chloride solution into a 600-ml, beaker. To the warm filtrate, carefully add distilled water heated to 60° C. until the solution just becomes



FIG 58 EDESTIN

turbid (300 to 500 ml. of water are required, depending upon the concentration of the protein). Warm the solution in a water bath at 60° until it becomes clear and then permit both the solution and the water bath to cool spontaneously at room temperature. In 24 hours there settles out a precipitate of globulin which is almost entirely crystalline in form. This particular globulin in hemp seed is called edestin. It is soluble in warm 10 per cent sodium chioride solution and may be crystallized by cooling its solution, or by dialyzing the solution against distilled water. Addition of warm water, as above, increases the yield of crystals obtained by decreasing the solubility of the protein. Edestin crystallizes in several different forms, chiefly octahedra (see Fig. 58). Filter off the edestin and make the following tests on the crystalline material.

Tests on Crystollized Edestin.

- Microscopicol Exomination (see Fig. 58).
- Solubility. Try the solubility in water, 1 per cent sodium chloride solution, dliute acid and alkali, and alcohol.
 - 3. Millon's Reaction.

4 Coagulation Test. Place a small amount of the globulin in a test tube, add a little water, and boil. Now add hydrochloric acid and note that the protein no longer dissolves It has been coagulated.

Dissolve the remainder of the edestin in about 50 ml. of 10 per cent sodium chloride solution, and make the following tests on this solution.

Tests on Edestin Solution.

 Influence of Protein Precipitants. Try a few protein precipitants such as nitric acid, tannic acid, picric acid, and mercuric chloride. Compare with the results on egg albumin (p. 174). Can you distinguish between albumin and globulin by tests such as these?

2. Biuret Test.

- 3. Coagulation Test. Boll some of the solution in a test tube. What happens?
- 4. Saturation with Sodium Chloride, Saturate about 18 ml, of the solution with solid sodium chloride, How does this result differ from that obtained upon saturating egg albumin solution with solid sodium chloride?
- 5. Precipitation by Dilution. Add a few drops of the solution to a test tube filled with distilled water. Why does the globulin precipitate?
- 6. Preceptation by Dialysis. Place the remainder of the solution in a dialyzing bag and dialyze against about 500 ml. of distilled water. Why does the globulin precipitate? Examine the precipitate under the microscope.

C GLUTELINS

It has been repeatedly shown, particularly by Osborne, that after extracting the seeds of cereals with water, neutral salt solution, and strong alcohol, there still remains a residue which contains protein material which may be extracted by very dilute acid or alkalt. These probably denatured proteins which are insoluble in all neutral solvents, but readily soluble in very dilute acids and alkalies, are called glutelins. The only member of the group which has yet received a name is the glutenin of wheat, a maxture of proteins which constitutes nearly 50 per cent of the gluten, the remainder being principally gladin.

Gluten: Preparation and Tests To about 50 g of wheat flour in a casserole or evaporating dish add a little water and mix thoroughly until a stiff dough results. Knead this dough thoroughly and permit it to stand for about half an hour. This is done in order that the maximum quantity of gluten may be obtained. Treat the dough with about 200 mi of water and knead it thoroughly. Note the yellowish color of the dough and the milky appearanced on a slide, cover with a cover slip, run underneath the slip a drop of lodine solution and observe the stained starch granules under the microscope?

[•] Work by Blish and San Iste it indicates that glutenin prepared by extraction of gluten with didute alkali is actually an irreversible alteration product of a more complex protein in the cruds gluten

Filter and apply a protein color reaction (see p. 169) to the filtrate. It should be positive, indicating that water-soluble proteins were present in the flour. Add fresh water to the dough and repeat the kneading process. Continue this procedure with fresh addition of water, or holding the gluten under the tap, until practically no starch granules are noted in suspension. To a small piece of the yellow, fibrous gluten apply Millon's reaction (p. 169). This test shows gluten to he protein material. Utilize the remainder of the gluten in the preparation of gliadin below.

Glutenin: Preparation and Tests. (In the preparation of gliadin below it is customary to remove this prolamin from the crude gluten by extracting with 70 per cent aicohol. Inasmuch as gluten consists chiefly of gliadin and glutenin the portion of the gluten remaining after the extraction of the alcohol-soluble protein gliadin may be utilized for the preparation of glutenin.)

To the finely divided residue from the preparation of gliadin below in a flask or hottle add ahout 250 ml. of 70 per cent alcohol. Allow to stand for ahout 48 hours with repeated shaking in order to remove any remaining gliadin. Crude glutenin remains. To purify the glutenin, treat it in a mortar with sufficient 0.2 per cent NaOH to dissolve it, and fliter the liquid through a wet pleated filter. Neutralize the filtrate earefully with 0.2 per cent HCi, adding the acid drop hy drop with thorough mixing after each addition. (The glutenin is soluble in excess of acid.) Filter off the glutenin precipitate and wash several times with 70 per cent alcohol and finally with water. Apply the following tests:

1. Solubility in water, 0.9 per cent NaCl solution, 0.2 per cent HCl, and 0.5 per cent NaCO.

2. Millon's Reaction.

D. PROLAMINS (ALCOHOL-SOLUBLE PROTEINS)

The term prolamin was proposed by Osborne for the group of plant proteins formerly termed "alcohol-soluble proteins." The name is very appropriate masmuch as these proteins yield, upon hydrolysis, especially large amounts of proline and ammonia. The prolamins are simple proteins which are insoluble in water, absolute alcohol, and other neutral solvents, but are soluble in 70 to 80 per cent alcohol and in dilute acids and alkalies. They occur widely distributed, particularly in the vegetable kingdom, and include zein of maize, hordein of barley, gliadin of wheat and rye, bynin of malt, and kafirin of kafir. Sorghum and rice also contain alcobol-soluble proteins. They yield relatively large amounts of glutamic acid on hydrolysis but little or no lysine. The largest percentage of glutamic acid (47 per cent) ever obtained as a decomposition product of a protein substance was obtained from the hydrolysis of the prolamin gliadin. This yield of glutamie acid is also the largest amount of any single decomposition product yet obtained from any protein except the 88.4 per cent of arginine obtained from salmin

Gliadin: Preparation and Tests. Introduce the finely divided crude giuten as prepared on p. 190 into a flask or bottle, add about 250 mi. of 50 to 70 per cent alcohol, and allow the mixture to stand 24 hours with occasional shaking. Filter (retaining the undissolved portion for preparation of giutenin, above) and evaporate the filtrate to dryness in a porcelain dish over a water

bath Pulverize the dry material Apply the following tests to this gliadin powder

Solubility and Protein Tests Test the solubility in alcohol (30 per cent, 50 per cent, and 70 per cent), water, 0 9 per cent \aCl, 0 2 per cent 11Cl, and 0 5 per cent Na:CO. Shake each tube repeatedly and filter To the filtrate apply the coagulation test (p 190) and the biuret test (p 171)

T. ALBUMINOUS (SCLEROI ROTLING)

The albuminoids yield hydrolytic products similar to those obtained from the other simple proteins already considered, thus indicating that they possess essentially the same chemical structure. They differ from all other proteins, whether simple conjugated, or derived in that they are insoluble in all neutral solvents. The albuminoids include "the principal organic constituents of the skeletal structure of animals as well as their external covering and its appendages" Some of the principal albuminoids are keratins from hair, horn, nails feathers, etc , elastins from arteries, ligaments, collagens from bone skin hoof, etc. reticulin, spongin, and silk fibroin Gelatin cannot be classed as an albuminoid al though it is a transformation product of collagen The various albuminoids differ from each other in certain fundamental characteristics (see p 183) Experiments on albuminoids will be found in Chapter 9

II CONJUGATED PROTEINS

Conjugated proteins consist of a protein molecule united to some non protein molecule or molecules otherwise than as a salt Glycoproteins nucleoproteins, chromoproteins phosphoproteins lipoproteins, and metalloproteins are the six classes of conjugated proteins

Glycoproteins may be considered as compounds of the protein molecule with a substance or substances containing a carbohydrate group other than a nuclcie acid. They yield upon decomposition protein and carbohydrate derivatives notably glucosamine CH₂OH (CHOHI)r CH(\H2) CHO galactosamine galactose fucose and uronic acids The principal Liycoproteins are (1) mucoids (2) mucins and (3) sulfomucins The elementary composition of typical mucoids is as follows

	N	s	c	И	0
Tendomucoid	11 7 ₉	2 33	48 76	6 53	30 60
Osscoinucoid	12 22	2 32	47 43	6 63	31 40

(For the preparation of tendomicoid see Chapter 9 of salivary much, see Chapter 13) Amyloid 10 which appears pathologically in the spicen, liver and kidneys, is also a glycoprotein

is not to be confused with the sulatance amyland which may be formed from cellulose (see p 20

Phosphoproteins are considered to be compounds of the protein molecule and phosphoric acid. The percentage of phosphorus in phosphoproteins is very similar to that in nucleoproteius, but they differ from this latter class of proteins in that they do not yield any purine bases upon hydrolytic cleavage. Two of the common phosphoproteins are the easein of milk and the ovovitellin of egg yolk. The phosphorus in these proteins exists in phosphorie acid radicals bound in ester linkage to the hydroxy amino acids serine and threonine. For the preparation of a typical phosphoprotein (casein), see Chapter 8.

Chromoproteins are compounds of the protein molecule with a metalloporphyrin (see Chapter 22) or some similar substance. The principal members of the group are the hemoglobiu of the blood and the chlorophyll proteins of plants. Upon hydrolytic cleavage, hemoglobin yields a protein termed globin and a coloring matter which contains iron and is known as heme. Hemoeyanin, another member of the class of chromoproteins, occurs in the blood of certain invertehrates, notably cephalopods, gasteropods, and crustacea. Hemocyanin generally contains either copper, manganese, or zinc, as contrasted to the iron of the hemoglobin molecule. For the preparation of hemoglobin in crystalline form.

see Chapter 22.

Lipoproteins consist of a protein molecule joined to lecithin, cholesterol, and other fatty substances. They have been comparatively little studied and may possibly be mixtures of protein and lipide material. These unions are not split by treatment with ether but are by alcohol. especially after heat denaturation of the protein.

Nucleoproteins. For consideration of nucleoproteius, see Chapter 7. Metalloproteins. Numerous protein enzymes contain metals as an inherent portion of their molecules, eg., carbonic anhydrase is a zinc protein complex, arginase contains manganese or magnesium, tyrosiuase

contains copper, etc.

III. DERIVED PROTEINS

A PRIMARY PROTEIN DERIVATIVES

1. PROTEANS

Proteans are those insoluble, denatured protein substances which are produced from proteins originally soluble through the incipient action of water, enzymes, very dilute acids, etc. According to Osborne nearly all proteins may give rise to proteans, and the determining factor in the transformation is the hydrogen-ion concentration. The protean produced from the transformation of edestin is called edestan; that produced from myosin is called myosau, etc. The name protean was first given to this class of proteins by Osborne in 1900 in connection with his studies of edestin. It is but little used at present.

2. METAPROTEINS

The metaproteins are denatured proteins formed by the action of dilute acids and alkalies on some proteins, especially on the albumins and the globulins. The conversion of protein into metaprotein is accelerated by a

rise in temperature, taking place almost immediately at the boiling point. There is, at present, no definite evidence concerning the exact nature of the changes which the protein molecule undergoes during this transformation (See the section "Protein Denaturation and Its Reversal" in Chapter 5). The denatured protein behaves like a suspensoid, dissolving in dilute acids and alkales—ie when the particles are electrically charged—and flocculating when the solutions are brought to the isoelectric point. The suspensoid character of the metaproteins is also indicated by their sensitivity to electrolytes especially in the neighborhood of the isoelectric point. The metaproteins are precipitated by saturation of their solutions with ammonium sulfate and when dissolved in acid, by full saturation with sodium chloride Bohing an isoelectric suspension of metaprotein converts it into coagulated protein, in which case the material is no longer soluble in dilute acids and alkales.

The metaproteins are usually classified as (1) acid metaproteins, or so-called acid albuminates and (2) alkali metaproteins, or alkali albuminates depending upon whether they are formed by the action of acid or alkali on the native protein Transformation products with similar physical properties are formed by the action of enzymes ultraviolet light, etc., on native protein It is unlikely that these procedures ever produce a homogeneous metaprotein

EXPERIMENTS ON A DENATURED PROTEIN

Preparation and Study Alkali Metaprotein (Alkali Albuminate) Carefully separate the white from the yolk of a hen segg and place the former in an evaporating dish Add 10 per cent sodium hydroxide solution, drop by drop stirring continuously The mass gradually thickens and finally assumes the consistency of jelly This is solid alkali metaprotein or "Lieberkuhin's jelly Do not add an excess of sodium hydroxide or the jelly will dissolve Cut it into small pieces, place a cloth or wire gauze over the dish, and by means of run ning water, wash the pieces free from adherent alkali Now add a small amount of water, which forms a weak alkaline solution with the alkali within the pieces, and dissolve the jelly by genthe heat Cool the solution and divide it into two parts Proceed as follows with the first part Neutralize with dilute hydrochioric acid, noting the odder of the liberated hydrochioric acid, noting the odder of the liberated hydrochioric acid, noting the odder of the liberated hydrochioric axid, ydrochioric axid hydrochioric axid hydrochioric axid hydrochioric axid hydrochior

- I Solubility Solubility in water, sodium chloride solution, dilute acid, and alkali
 - 2. Millan's Reaction
- 3 Congulation Test Suspend a little of the metaprotein in water (neutral solution) and heat to boiling for a few moments Now add I to 2 drops of di lute NaOII solution to the water and see if the metaprotein is still soluble in dilute alkali. What is the result and why?
 - 4 Test for Cystine and Cyste ne Sulfur (see p 168)

Subject the second part of the original solution to the following tests

5 Congulation Test Heat some of the solution to boiling in a test tube

6. Biuret Test.

 Influence of Protein Precipitants. Try a few protein precipitants such as pictic acid and mercuric chloride. How do the results obtained compare with those from the experiments on egg albumin? (See p. 174)

3 COAGULATED PROTEINS

Simple proteins, such as the albumins and globulins, are converted by heat, ultraviolet light, mechanical agitation, or long contact with alcohol, etc., into insoluble materials known as coagulated proteins, which probably consist of the matting together of the denatured protein fibrils similar to the formation of felt from short fibers of hair (See p. 156 and the section "Protein Denaturation and Its Reversal" in Chapter 5.) Coagulated proteins are insoluble in water, salt solutions, and dilute acids and alkalies. They are soluble in strong milieral acids and alkalies which hydrolyze the protein into simpler substances.

EXPERIMENTS ON COAGULATED PROTEIN

Ordinary coagulated egg white may be used in the following tests:

 Solubility. Try the solubility of small pieces of coagulated protein in water, 1 per cent sodium chloride, and dilute and concentrated acid and all all

2. Millon's Reaction

- 3. Xonthoproteic Reaction Partly dissolve a medium-sized piece of the protein in concentrated nitric acid. Cool the solution and carefully add an excess of ammonium hydroxide. Both the protein solution and the undissolved protein will be colored orange.
- 4. Bruret Test. Partly dissolve a medium-sized piece of the protein in concentrated sodium hydroxide solution. If the proper dilution of copper sulfate solution is now added, the white coagulated protein, as well as the protein solution, will assume the characteristic purplish-violet color.

B SECONDARY PROTEIN DERIVATIVES

1 PROTEOSES AND PEPTONES

The proteoses and peptones are poorly defined intermediate products formed during the digestion of proteins by the proteolytic enzymes, as well as in the decomposition of proteins by hydrolysis and the putrefaction of proteins through the action of bacteria. As hydrolysis proceeds, the large, colloidal protein molecule is split up into a mixture of large and small fragments which were formerly designated as primary and secondary proteoses, peptones, and peptides. The larger fragments are constantly being broken down until, finally, only amino acids remain. It should be emphasized that the substances formerly known as proteoses and peptones are not homogeneous chamical entities, but rather mixtures of frigments of the original protein molecule which vary in composition and also in size, and ire grouped together merely because they exhibit

certain characteristic properties in common. Those fragments which are precipitated when the solution is half saturated with ammonium sulfate were known as primary protoces, those precipitated when the solution is saturated with ammonium sulfate were called secondary protocess. The peptones were not precipitated by ammonium sulfate. The protocess still exhibit colloidal properties, but these properties become less pronounced and begin to disappear entirely by the time the breakdown of the protein molecule has reached the peptone stage. Thus, in addition to being nonprecipitable by ammonium sulfate, the peptones diffused through such membranes as collodion and failed to give some of the obaracteristic precipitation reactions of roteins.

Since the proteoses and peptones are both heterogeneous mixtures of fragments of the original protein molecules different preparations of these substances will show variations in composition and properties depending upon the nature of the protein used as starting material, the method of hydrolysis employed, and the method used to separate the various products of hydrolysis Thus the so-called peptones sold commercially vary not only in composition but also in the amounts of primary and secondary proteoses they contain As a class the proteoses and peptones are a mixture of amino acids and peptides not coagulated by heat Some are also soluble in dilute alcohol Peptones differ from proteoses in being more diffusible, and nonprecipitable by (NII4)2804, and by their failure to gue any reaction with potassium ferroe janide and acetic acid, potassio-mercuric todide and HCl, pieric acid, and trichloroacetic acid Peptones may be precipitated by phosphotungstie acid, phosphomolybdic acid absolute ilcohol, and tannic acid, but an excess of the precipitant may dissolve the precipitate. The so-called primary proteoses, being relatively large molecules, are precipitated by HNO, and are the only members of the proteose-peptone group which are so precipitated

EXPERIMENTS ON PROTEOSES AND PEPTONES

Some of the more general characteristics of the proteose peptone group may be noted by making the following simple tests on a proteose peptone powder

- I Solubility Solubility in hot and cold water and sodium chloride solu-
 - 2. Millon s Reaction
- 3 Precipitation by Picric Acid To 5 ml of proteose peptone solution in 2 test tube add picric acid until 2 permanent precipitate forms. The precipitate disappears on heating and returns on cooling
 - 4. Precipitation by a Mineral Acid Try the precipitation by nitric acid.
 - 5 Congulation Test lient a little proteone peptone solution to boiling bose it congulate like the other simple proteins studied?

SEPARATION OF PROTEOSES AND PEPTONES¹¹

Place 50 ml. of proteose-peptone solution in an evaporating dish or casserole, and half-saturate it with ammonium suifate solution, which may be accomplished by adding an equal volume of saturated ammonium suifate solution. At this point note the appearance of a precipitate of the primary proteoses. Now heat the half-saturated solution and its suspended precipitate to bolling and saturate the solution with solid ammonium suifate. At full saturation the secondary proteoses are precipitated. The peptones remain in solution.

Proceed as follows with the precipitate of proteoses: Collect the sticky precipitate on a rubber-tipped stirring rod or remove it by means of a watch glass to a small evaporating dish and dissolve it in a little water. To remove the ammonium sulfate, which adhered to the precipitate and is now in solution, add barium carbonate, boil, and filter off the precipitate of barium sulfate. Concentrate the proteose solution to a small volume¹³ and make the following tests:

Tests on Proteoses.

- 1. Biuret Test.
- 2. Precipitation by Nitric Acid. What would a precipitate at this point indicate?
- Precipitation by Trichloroccetic Acid. Tbls precipitate dissolves on heating and returns on cooling.
 - Precipitation by Picric Acid. This precipitate also disappears on heating and returns on cooling.
 - 5. Precipitation by Patassio-mercuric Iodide and Hydrochloric Acid.
 - 6. Coogulotion Test. Boll a little in a test tube. Does it coagulate?
 - 7. Precipitation by Acetic Acid and Potossium Ferrocyonide.

Tests on Peptones. The solution containing the peptones should be cooled and filtered, and the ammonium sulfate in solution removed by boiling with barlum carbonate as described above. After filtering off the barlum sulfate precipitate in the presence of a filter aid such as diatomaceous earth, concentrate the peptone filtrate to a small volume and repeat the tests as giveo under the protose solution, above. Also try the precipitation by tannic acid. In the bluret test the solution should be made very stroogly alkaline with solid potassium hydroxide.

¹¹ The separation of proteoses and peptones by means of fractional precipitation with ammonium sulfate does not possess the significance it was once supposed to possess insening the boundary between these substances and peptidesis not well defined (see p. 113) for discussion of a quantitative method for determining protein and proteose based on their separation by trichlorosectie and see Seibert J. Biol. Chem. 70, 205 (1926).

If the proteoses are desired in powder form, this concentrated proteose solution may now be precipitated by alcohol or acctone, and this precipitate, after being washed with acctone and with ether, may be dired and powdered.

25

Intrato may contain econdary cloteser (Tr.) nay tests given on p 347.)

Precipitate Indicates pelatu (1ry ans gelatin

Hitrate Stitutote with a sted (NHO)-No. Irectificate fare librances according to professor of adjusten Hit-tree It off and dissolve in

I reclifiate full enter planden primity 110 fease or gelatin (Ity tertosa color leats)

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Schulb for Derector of Profess

lent to lading sorm tealer. bergente al d ohn 15 pancing some of the unknown solution into 29.1 i 40 adduces of water Precis Itato tada atos globules. Net recij unte indenten premary refessa. or golden " ----

No congulation in kenten second my positions (Lry protein color text) Centralistica lecti-

o Galatu and retecous and be deferentated 15 no cas of the 110 Line Cole rarches (see p. 170). A rould'us reaction here would fusion to retrose and a matrix o reservance NOB Relatin.

MODEL CHART FOR REVIEW PURPOSES

		2	Solul	pyrt	,		Preceptation Tests						ang ut ata					
Protein	ll ater	10 ", NaCi	020, 1101	OS Na,CO,	Co t HCt	Cone NOII	I rotein Colar Test	Vi eral dead (HNO)	Metallic Salt (HgCh)	Alcohol	I of Ferrocvanide	I oldano-n ercursc Io I da + IICl	I teric Acid	Tricilorouceise Acid	(NII4)2504	NaCl	Difft ston	Coagulatio by Heal
Album p				1		}	1				}							
Globulin			1	1	1	1												
\ucleoprate n	1		-							-								
Phosphoprote n	1	1	1		1	1			1									
Gl; coprote a		1		1		1			_	Γ			-	Γ		_		
As d metaprote n			Ī	1					-									
Alkalı metaprote u	Γ	-					ĺ				1							
Proteose					1	-	1	1		j	1							
Peptone							T	1		1								ì
Coagulated prote n		T	T	1	1	T	1	Γ		Ī						_		

2 PEPTIDES

The peptides are "definitely characterized combinations of two or more amino ands, the carboxyl (COOH) group of one being united with the amino (NH-) group of the other with the elimination of a molecule of water" These peptides are more fully discussed on pp 113 and 152

REVIEW OF PROTEINS

In order to facilitate the student's review of the proteins, the preparation of a chirt similar to the model shown above is recommended. The signs + and - may be conveniently used to indicate positive and negative reactions.

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(5 methyl-uracil)

With all these structures, tautomerism from lactim to lactam forms is possible, for example

Tautomeric forms of uracil

The Nucleotides and Nucleosides. Each purms or pyrimidine in a auctive acid is found to be accompanied by one molecule of pentose and one of phosphoric acid. I he studies of Levene, Jones, and others demonstrated that hydrolytic cleavage of the nucleus acids could lead to phosphoribosides, or nucleotides of each of the bases. The nucleus acids are polymers of minip such nucleotides and hence the descriptive name polynical obstacles as auxiliation in further hydrolysis is involved, the phosphoric acid is removed and the ribase derivatives of the bases, the nucleosides, are obtained. The relation between nucleosides and nucleotides may be indicated in the diagram and table shown on p. 203

The nucleic ands used most extensively in the studies of their chemistry have been the ribose nucleic and from yeast or yeast nucleic acid, and the decryphose nucleic acid from thymus gland, or thymus nucleic acid. They contain approximately equimolar amounts of each of four

bases This led to a long prevalent theory that the nucleic acids were tetranucleotides, or at least contained a repeating tetranucleotide unit. However, with the advent of more refined techniques, analyses now indicate that few nucleic acids contain the equimolar quantities of the four bases required by the tetranucleotide theory. The presence of a fifth nucleotide in some deoxymbose nucleic acids also discredits that hypothesis. Present concepts of nucleic acid structure suggest a much greater complexity. These large molecules are still less well understood than are the proteins.

	Nucleatide	
Purme or	Ribose or	Phosphoric
Pyrimidino	Deoxyribose	Acid
Nucl	enside	

RIBOSE NUCLEIC ACID DERIVATIVES

Base	\ ucleoside	Nucleotide
Uracil	Uridine	Undylic Acid
Cytosine	Cytidine	Cytidylic Acid
Adenine	Adenosino	Adenylic Acid
Guanme	Guanosine	Guanylic Acid

DEOXYRIBOSE AUCLEIC ACID DERIVATIVES

Thymine Thymidine Thymidylic Acid
For the deoxyribose derivatives of the other three bases
the prefix deoxy is used with the names above, as in
deoxycylidine

Purine and Pyrimidine Content. Analyses of the purine and pyrimidine content of various deoxyribose nucleic acids by paper-coromatographic methods have shown variations in the composition of the DNA from various species Chargaff has shown that though the ratio of total purnes to total pyrimidines is usually close to unity, there are types in which the amounts of one purme and one pyrimidine may considerably exceed the amounts of the others It is of interest that, of the various possible combinations only two types have been encountered one in which adenue and thymine predominate, and one in which guanine and cytosine predominate The former is most frequently found in mammalian tissues, yeast and so forth, the latter is common in bacteria. Some bacteria possess a DNA with nearly equal proportions of the four bases. Where 5-methyl cytosme is found to be present as a fifth component (in plant. fowl, and mammalian species, including man) it is usually found only in traces, although in the case of the DNA of wheat embryo it amounts to oue-eighth of the total base present

The Amount of Nucleic Acids in the Cells.

(a) Decyamose Nucleic Acts The total quantity of decymbose nuclea and per cell is remarkably constant in all the normal sometic inssues within a given species. This amount is not altered by conditions such as sturvation or other stress. The amount per hiploid (sperm) nu

cleus is one balf of that for a normal diploid nucleus of the species and where polyploidy is found the DNA per nucleus is correspondingly in creased. There are considerable variations from species to species fish and birds having generally a smaller amount of DNA per nucleus than mammals (which generally contain in the order of 6 × 10 ° mg per nucleus). The constancy of the amount of DNA per nucleus and of the composition throughout a species correlates well with the requirement of the genetic theory for a constancy in the character of the chromosome. This is another of the lines of evidence which foster the theory that DNA may be an integral component of the chromosome.

(b) RIBOSE VICLEIC ACID The ribose nucleic acids of various tissues vary in composition from tissue to tissue and that of the nucleus differs from that of the cytoplasm Preparations of RNA satisfactory for ana lytical studies are few and no generalizations can be made. The quantity of RNA per cell fluctuates under various conditions. It is greatly in fluenced by such factors as the nutritional state of the animal and is greater in those cells with higher metabolic activities. For instance it is bigh in secretory organs such as salivary glands and pancreas where active protein synthesis and secretion are taking place. There is also relatively more RNA in rapidly growing embryonic and tumor tissues Much circumstantial evidence is available which indicates that there is an increase in quantity and metabolic activity of the RNA just prior to or associated with protein synthesis and the theory that ribose nucleie acid plays a major role in the synthesis of protein has received much support However there is yet no concrete evidence of any direct participa tion of RNA in protein synthesis Several other suggestions that RNA may be primarily involved in processes such as secretion or energy me tabolism have some support. It remains that ribose nucleic acid is a dynamic component of the cell and is certainly associated with active metabolic processes but a definite assignment of a functional role is not yet possible

THE STRUCTURES OF NUCLEOTIDES AND OF POLYNUCLEOTIDES

The pentoses in the nucleosides are attached in a nonreducing N gly cosyl linkage. Note that the formation of a pyrimidine nucleoside requires that the 1 and 2 positions of the pyrimidine be in the lactam form. The positions of attachment, the existence in the furances form and the ster cochimical configurations have been fully authenticated. In the case of the ribose nucleotides, three isomers are possible depending upon which one of the hydroxyl groups of the sugar is esterified with the phosphore acid. In the case of adenylie acid all of the three possible isomers are known (see p. 205)

Two isomers of each of the purme or pyrimidine nucleotides are produced upon alkaline hydrolysis of ribose nucleuc ands. These are the 2 and 3 isomers which were also known as the a and b isomers during a long uncertainty as to which was which. The chemical properties of the isomerse nucleotides are very smilar but it is possible to separate them by ion-exchange chromatographic te him pues. The name. jeast aden, jile

acid formerly referred to such a mixture of these isomeric adenylic acids, as obtained from yeast nucleic acid. The discovery that there were not four nucleotides, but four pairs of isomeric nucleotides that could be obtained from ribose nucleic acids, necessitated re-evaluation of many of the structural and metabolic aspects of the polynucleotides.

One of the isomeric Adenylic Acids (9-β-p-ribofuranosyl-adenine-3'-phosphate) (Adenosine-3'-phosphate)

One of the isomeric Uridylic Acids (1-β-n-nbofuranosyl-uracıl-2'-phosphate) (Uridine-2'-phosphate)

The third possible isomer of adenylic acid is the one in which the phosphate is attached at the 5' position. This nucleotide, adenosine-5'-phosphate, has long been known to occur in muscle and bears the name nuscle adenylic acid. It is the adenylic acid from which adenosinetriphosphate (ATP) and certain coenzymes are derived, and it thus plays important roles in carbohydrate metabolism, fermentation, and muscular contraction (see Chapter 10).

Enzymatic degradation has provided further information on the nature of the linkages present in RNA. Cleavage of RNA by pancreatic ribonuclease liberates the majority of the pyrimidines as the 3'-cytidylic and 3'-uridylic acids. An enzyme-resistant portion remains which is composed chiefly of purine nucleotides. The subsequent action of a phosphodiesterase (prepared from snake venom) leads to the production of all four 5'-mononucleotides, chiefly purine nucleotides but also small amounts of pyrimidine nucleotides. These results suggest that it is the 3' and 5' linkages which predominate in the nucleic acid, despite the fact that it is the 3' and 2' isomers that are produced by the action of alkali

The production of the mixture of the 2' and 3' isomers upon alkaline hydrolysis of nucleic acids is attributed to the formation of cyclic phosphate esters, such as that depicted here. In this illustration the phose and its three available hydroxyl groups are presented schematically.

position of first cleavage by alkali

In this treater of phosphoric acid the bond which is not involved in the cyclic phosphate is the most labile to alkali and is cleaved first. The cyclic mononnel cotides which are thus liberated are then further cleaved by alkali to yield mixtures of the 2' and 3' mononnel cotides.

In the deoxyribose nucleic acids the lack of a hydroxyl in the 2' position of the sugar prevents the formation of the cyclic phosphate, as 18 evident from the formula of thymidine shown on p 207 The deoxyribose nucleic acids are thus more resistant to depolymentation by alkali, in fact, most methods for separation of the two types of nucleic acid depend upon the difference in their stability in the presence of alkali.

In the ribose nucleic acid the presence of approximately one ionizable hydrogen per phosphate indicates that most of the phosphates are present as diesters Approximately one quarter of this titratable hydrogen shows a pK of about 6 corresponding to a secondary phosphoryl dissociation, and this indicates that the ribose polynicietudes are not solely diphosphate ester bridges from the 3 to the 5' hydroxyls of adjacent nucleotides beviral other lines of evidence suggest the presence of some monosterified ribose (i.e., oxidation by periodic acid production of a dimethyl n-bost after methylation), of some triesterified ribose (production of free

ribose after methylation), and of some triesterified phosphoric acid (from titration data). The characterization of a number of dt-, tri- and tetra-nucleotides, which are obtained by partial enzymatic or acid hydrolyses of ribonucleic acids, also reveals that several purine or pyrimidine nucleotides may be adjacent to one another. Sufficient information is not available to plot the sequence of nucleotides in any nucleic acid.

A formulation such as that shown in Fig 59 is compatible with the present chemical and enzymatic evidence and indicates the complexity of the structure of the polynucleotides known as ribose nucleic acids. This

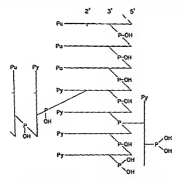


Fig. 59 Fragment of a Ribose Nicleic Acid Schematic presentation of the types of linkage myolved. (Atter Coha, Dobetty, and Volkan Phosphorus Metabolism) dol. 2 Baltunore, Johns Hopkan University Press, 1952.

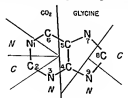
sketch of the current speculations regarding intermediate linkages represents only a portion of a molecule A molecular weight of 30,000, for instance, requires nearly 100 nucleotides

Vanous enzymatic hydrolyses of the deoxyrihose nucleic acids have yielded 3' nucleotides or 5' nucleotides Small amounts of deoxyrihose nu cleosides bearing two phosphore acids (in both the 3' and 5' positions) have also heen obtained. The nucleotides in DNA must be joined prin cipally by 3', o' phosphate dester hidges, with possibly some hranching through triesterfied phosphate.

It should be pointed out that phosphate bridges between the 5' hydroxyls of two nucleotides have not been found, nor have pyrophosphate linkages been implicated The latter type of linkage is found in those coenzymes which are formed from adenosine 5' phosphate and various stamin derivatives. Despite their considerable similarities there is thus a fundamental difference between nucleoproteins and those enzymes which are complexes of a disucleotide coenzyme and a protein (see Chapter 12)

METABOLISM OF NUCLEIC ACIDS

The organism is able to synthesize de note all of its nucleic acids and none of the organic constituents thereof are necessary in the diet. It has therefore here difficult to learn what compounds are utilized in forming the tissue nucleic acids, since classical nutritional and halance studies were able to continute tittle. Viost of the knowledge of the origin of the nucleic acid components has come from the administration of isotopically labeled compounds and the use of degradative procedures which permit analysis of individual moieties of the products. In this way, the principle sources of the individual atoms of the purmer ring have been shown to be



FI 60 ORIGIN OF THE ATOMS OF THE PURINE RING

C from 1-carbon sources (formate formaldehyde) V—from VH; (via aspariate glutamate glutamine) From B own, Roll, and Wenfeld Fassphorus Medicales, Vol. 2. Baltimore Johns Hopk as Layersity Press, 13 2.

The 4 rearbon and 7 nitrogen portion is derived from glytine as a single unit, the 6 carbon arises from carbon dioxide. The 2 and 8 carbons arises.

from one-carbon units such as formate or formaldehyde, and folic acid plays a direct role in the incorporation of these one-carbon units. There is no evidence that there is any formation of free purines in the course of biosynthesis in tissues. Rather, the ribose and phosphate are attached to a smaller precursor before the synthesis of the purine ring is fully completed. This has been shown to be true in studies of inosinic acid (bypovanthme ribotide) biosynthesis in pigeon liver, and the mechanism of synthesis of purine derivatives in this species is probably identical or very similar to such synthesis in the manimal.

In birds and reptiles there is a peculiarity of purine metabolism not observed in mammals. In those species ure acid, instead of urea, is formed directly as the chief catabolic end product of protein introgen metabolism.

Among the purmes classically associated with purme metabolism, only adenine reaches the nucleic acids in significant amounts when fed to the rat. Adenine is not only incorporated as adenine but is also partially transformed into the guanne of the polynucleotides, and by the use of isotopes it has been demonstrated that the purme ring remains intact during this conversion. Guanne and several other purmes are extensively catabolized and, in the rat do not reach the polynucleotides. Guinne ean, however be readily used by several species of microorganisms and to a small extent by the mouse, and in most cases it can also be transformed into polynucleotide adenine. A purme which has not been shown to occur in nature, 2,6 diaminopurine (or 2 aminoadenine), can be utilized for the synthesis of polynucleotide purnes. In the rat it is converted only into nucleic acid guanne although in some species it may be converted to both nucleic acid guanne.

The nucleoside and nucleotides of adenine may be utilized by the rat but only if they are administered parenterally in order to avoid the diges tive enzymes of the intestinal tract Guanylic acid can lead to nucleic acid guaniue, although guanosine like guanine fails to do so in the rat

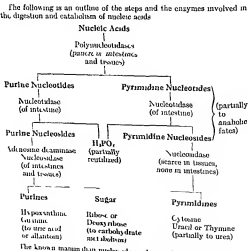
Less is known of the origin of pyrimidines. The 2 carbon can arise from carbon dioxide, and aspartie acid and carbamyl aspartic acid are major precursors of the other carbons of the ring. The pyrimidines uracil and cytosine are not significantly utilized for polynucleotide synthesis, al though their nucleosides or nucleotides are very effectively utilized. There is an interconversion of the pyrimidine derivatives, similar to that of the purmes, and thus cytidine and cytidylic acid are transformed into ribose nucleic acid uracil and also into deoxyribose nucleie acid cytosine and thymine The conversion to the latter involves the acquisition of an addi tional carbon for the 5-methyl group of the thymine This arises from one-carbon units such as formic acid or the \$ earbon of serine, and folic acid is also involved in this reaction. Vitamin B12 is likewise involved in the formation of thymidine and other deory ribosides but its role is not yet adequately defined It is of interest that a pyrimidine which is found in milk profit acid or 6-carboxyuracil is readily utilized as a precursor of polynucleotide pyrimidines. Its possible significance in infant nutrition is

The usual diet contains few sources of individual purines, pyrimidines

and their nucleosides or nucleotides Varying amounts, however, of nu cleic acids will normally be encountered in the diet particularly when appreciable quantities of glandular meats are consumed. The feeding of isotopically labeled ribose (yeast) nucleic acid to the rat showed that there was appreciable incorporation of its pyrimidines, but no significant incorporation of its purines into the tissue nucleic acids. This is in distimet contrast to the fact that when the individual free pyrimidines are present in the diet they are not incorporated. Since adenine and its derivatives can be incorporated it is obvious that the purine moieties of the nu clue and are catabolized further than the nucleotide or nucleoside stage, and that the catabolic products do not include adenine On the other hand the pyrimidine moieties which were utilized must have been absorbed before degradation to the free pyrimidines. The fate of the dictary nucleic acid can be correlated with the known fates of the various derivatives and with the known degradative enzymes described below

ENZYMATIC BREAKDOWN OF NUCLEIC ACIDS

the digestion and catabolism of nucleic acids



The known mamm dun nucleosidases do not cleave adenosine but do split most is (hypoxanthine ribeside) and guanosine. The very active adenosine deaminase does, however, convert adenosine to inosine, and the net result is to lead to purines which are extensively catabolized and not to purines which are utilized for nucleic acid synthesis. The lesser abundance of pyrimidine nucleosidases does not result in the rapid destruction of the ribose derivatives of the pyrimidines.

CATABOLISM OF PURINES

The purines are readily catabolized to uric acid via a series of deaminations and oxidations as outlined below. Adenosine and its derivatives can reach hypoxanthine through deamination to inosine and cleavage of the inosine to hypoxanthine If a large amount of adenine is present it can be oxidized directly to 2,8-dioxyadenine, which may cause kidney damage through deposition of crystals of this very insoluble substance in the tubules.

In man and apes, uric acid is the primary end product of purine metabolism, but in other mammals an enzyme, uricase, carries the oxidation

further to the more soluble allantoin. In man intrivenously administered uric acid is almost completely exercted as such and only a small but variable amount is degraded as far as urea. However, orally administered uric acid is extensively degraded to urea, presumably by intestinal bacteria

In gout the blood une acid concentrations are increased somewhat above normal, and the ultimate result of the disease is the deposition of toplu of sodium urate in the joints and clsewhere. However the uric acid clearance is usually essentially normal although it may be decreased where renal impairment has resulted

In the normal individual there is somewhat less than a gram of unc acid in the blood and fluids in a volume approximately equal to the extracellular space The size of this available pool, and its rate of renewal, can be measured by the intravenous injection of a small sample of isotopic uric acid and determination of the course of excretion of the isotope Benedict and Stettin have shown that in the gouty individual the size of the pool may be increased many fold Only the outer portions of the tophaceous deposits of urates were found to equilibrate rapidly with ad munistered isotopically labeled urie acid. The chology of gout is complex but a study making use of isotopic glycine has confirmed the view that increased synthesis of une acid is a factor

The administration of ACTH leads to uncaciduria, and in the gouty patient can precipitate an attack of acute gouty artbritis Increased blood uric acid is one of the outstanding changes observed in eclampsia

EXPERIMENTS ON NUCLEOPROTEINS AND THEIR DERIVATIVES

- I Preparation of Deoxyribosenucleoprotein from Thymus or Spleen Fresh tissue is minced and thoroughly washed with physiological saline, containing 0 01 M sodium citrate The washed tissue is then extracted by stirring with about 10 volumes of I M \aCl containing 0 01 M sodium citrate The suspen sion is centrifuged at high speed (10,000 to 12 000 r p m) and the supernatant ilquid containing the nucleoprotein is removed. This solution is poured into 6 volumes of water (which reduces the \aCl concentration to that of an 180 tonic solution), and the nucleoprotein precipitates as a fibrous mass. The supernatant liquid is decanted and the nucleoprotein is purified by again dissolving in WaCl The solution is centrifuged at high speed to remove any suspended material, and the nucleoprotein is again precipitated by pouring into 6 volumes of water If the mixture is stirred with a rod having a crook at lts end, the fibrous material generally winds around the rod and adheres to it
 - 2 Tests on Nucleoprotein Try the following tests on a nucleoprotein preparation
 - a Try the xanthoproteic Willon and hiuret tests
 - b Test the solubility in water, 10 per cent NaCl, 10 per cent HCl dilute kOII, and alcohol
 - c Test for Phosphorus in Organic Matter To a small amount of the sub stance in a crucible add about five times its bulk of fusion mixture (two parts

Mirsky and Iollater J Cen I April 30 117 (1916) Letermann and Lamb J Bil Chem. 1 6 65w (1345)

sodium carbonate to one potassium nitrate). Heat carefully until the resultlng mixture is coloriess. Cool, dissolve the mass in a little warm water, acidify with uitric acid, heat to about 65°C., and add a few ml. of molybdate solution. In the presence of phosphorus a yellow precipitate of ammonium phosphomolybdate is formed.

Instead of acidifying with nitric acid, the aqueous solution may be approximately neutralized with hydrochloric acid, a few mi. of magnesia mixture added, and then excess of ammonium hydroxide solution. A white precipitate of magnesium ammonium phosphate is formed.

- d. Dissolve a little of the nucleoprotein in very dilute KOH and then make slightly acid with acetic acid. Explain results.
- 3. To Show the Presence of Protein, Carbohydrate, Phosphoric Acid, and Purine Radicals in Nucleoprotein. Upon complete acid hydrolysis of nucleoprotein, these substances will be liberated, as well as decomposition products of the protein part of the molecule. To show their presence, proceed as follows: Transfer about a gram of nucleoprotein to a small flask and add 50 ml. of 5 per cent H2SO4. Boil for an hour or more to hydrolyze. Maintain the original volume by adding water. The solution becomes brown owing to the formation of melaninlike substances. Cool and filter the acid solution. Apply the following tests to portions of it: (a) the hiuret test, (b) the xanthoproteic test. (c) the Molisch test, (d) Benedict's test, and (e) the test for phosphate. (f) Transfer about 25 ml. of the hydrolyzate to a casserole and add ammonia with thorough mixing, a little at a time, until the fluid is nearly neutral. Then make slightly alkaline with dilute ammonia and filter if not clear, Transfer to a beaker and add about 10 ml. of 5 per cent ammoniacal silver nitrate solution. Purine bases will yield a flocculent precipitate of their aliver salts. If a precipitate does not appear immediately, examine the solution after it has been allowed to stand undisturbed for some time.
 - d. Preparation of Ribose Nucleic Acid from Yeast. Dilute 50 ml. of 1 per cent NaOII with 250 ml. of water in a casserole and add to this solution 100 g. of compressed yeast cut in small pieces or 30 g. of dry yeast. Heat on the water bath for half an hour with occasional stirring. Remove from the hath and filter at once through a folded filter. To the cooled filtrate add acetic acid until faintly acid to litmus. Filter again Evaporate the solution to 100 ml. or less, and filter if necessary. Allow to cool to 40°C. or below, then pour with vigorous stirring into 200 ml. of 95 per cent alcohol containing 2 ml. of concentrated HCl. Allow to settle in a tall vessel and decant. Wash twice with 95 per cent alcohol and twice with ether. Transfer to a filter paper. Allow to drain and dry at room temperature.
 - 5. Tests on Nucleic Acid from Yeast.3
 - a. Test the solubility of nucleic acid in cold and hot water, in alcohol, and in dilute acid and alkali. To the solution in alkali add dilute IICI drop by drop until the solution is acid; then add excess of concentrated IICI.

Does nucleic acid coaguiate on boiling? Does the solution in hot water geiatinize on cooling?

b. Try the bluret test.

t

c. Dissolve a little nucleic acid in water with the aid of heat. Test the reaction of different portions of the solution with litmus, alizarin, and Congo red solution.

A satisfactory preparation of yeast nucleic and may be obtained from Schwarz Laboratories, Inc., Mount Vernon New York

d Boil a small portion of the nucleic acid with about 10 ml of 10 per cent sulfuric acid for one to two minutes Divide into three portions

sulfuric acid for one to two minutes. Divide into three portions.

To one portion apply carbohydrate tests—e g, the a-naphthol (Mollsch) reaction and Tauber stest. What do these indicate?

To a second portion apply a test for purine hases. Add an excess of ammonia and then a little silver nitrate solution.

To the third portion apply the test for phosphate, adding ammonia in slight excess, then making acid with nitric acid, adding molybdate solution, and warming

6 Preparation of Deoxyriboie Aucleic Acid Freshly excised thymus glands are minced and frozen The subsequent extractions are carried out as near 6°C as possible Homogenize 100g of frozen tissue in 200 mil of coid 0 15 M NaCl solution, which is also 0 01 M in sodium citrate, and centrifuge the mixture at 2000 r p m This operation is repeated three times on the residue and the supermatants are discarded

The residue is dispersed in about 200 ml of water' in a high speed blendor Wincontinued stirring sufficient 5 per cent sodium dodecyl sulfate is added to bring the concentration to 0 5 per cent, and sufficient NaCl to make this solution i 'N with respect to it. The mixture is stored at 5'C overnight and then filtered through a pad of Celite on a Buchner funnel. When clear, the filtrate is poured, with stirring, into two volumes of ethanol. The fibrous precipitate of sodium deoxyribose nucleate is collected and is washed repeatedly with 70 to 59 per cent algobil.

7 Tests on Deoxyribosenucleic Acid Repeat Exps 5 (a) through (d) as given under yeast nucleic acid, above (e) Vaks a 4 per cent solution of deoxyribosenucleic acid in hot water (0.4 g. to 10 mi) Allow to cool What happens? Divide into two portions To one add a little NaOII solution, to the other add acetic acid Then neutralize carefully in each case

Both acetic acid and NaOH decrease the viscosity of the nucleate solution it may be changed back and forth from the gelatinous to the fluid condition by the alternate addition of acid and aikall

- 8 Tests on Purine and Parimidine Bases and Derivatives
 (a) Nanthine.
 - (i) Silver Nutrate Reaction Dissolve a little xanthine in ammonia and add silver nitrate solution Examine a little of the precipitate microscopically (see Fig. 61)
 - (2) Copper Sulfate Rection Dissolve a little of the substance in dilute alkali, make faintly acid with acetic acid Heat to boiling Add i mi of 10 per cent CuSD, and then a few drops at a time of sodium bisuffite (saturated solution) until the precipitate becomes yellowish All of the purines give this reaction.
 - (3) Vitre Acid Test Place a small amount of the substance in a small evaporating dish, add a few drops of concentrated nitric acid, and evaporate to dryness very carefully on a water bath The yellow residue upon moistening with caustic potash becomes red in color and upon further heating assumes a purplish red hue Now add a few drops of water and warm. A yellow solution results which yields a red repsidue.

³ For best results the solution should be analyzed for nitrogen and the nitrogen concentration adjusted to 0 o mg, per ml. Marko and Butler J Biol. Chem. 190 165 (1951)

upon evaporation. Compare with a similar reaction on other purine bases and uric acid. (See the murexide test, p. 796.)

- (b) Hypoxanthine.
 - (1) Repeat Exps. 1 and 3 under Xanthine. Examine the crystals of hypoxanthine silver nitrate under the microscope. (See Fig. 62.)

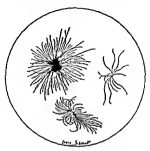


FIG 61 NANTHINE SILVER NITRATE.



Fig. 62. Hypoxanthine Silver Nitrate, Drawn from a student preparation by Dr E. F. Hirsch.

- (2) Dissolve a little of the substance in a very small amount of hot 6 per cent nitric acid and allow to cool. Characteristic whetstone crystals of hypoxanthine nitrate should be formed. Examine under the microscope.
- (c) ADENINE.
 - (1) Dissolve a little adenine in hot water and add a few drops of picric acid. Examine the pale yellow clusters of needles under the microscope.
 - (2) Repeat Exp. 3 under Xanthine.

(d) GUANINE

(1) Dissolve a little of the substance in 20 to 25 times its weight of hoiling 5 per cent alcohol. Allow to cool and examine the crystals under

the microscope (2) Dissolve a little guanine in 20 to 25 times lts weight of holling 5 per

cent hydrochloric acid Allow to cool, and examine the crystals.

(3) Perform Exp 3 under Xanthine

(e) Unic Acid On a small amount of uric acid, try Test 3 as given under Xanthine This test on uric acid is called the murexide test. For other tests on uric acid see Chapter 28, Urine.

(f) REACTIONS FOR URACIL CYTOSINE AND THYMINE Treat an aqueous solution of each pyrimidine with an excess of bromine. Remove the excess by boiling the solution. Add harium hydroxide in excess. A purple color indicates cytosine or uracil and is due to the purple barium salt of dialuric acid

Reflux for a few minutes and distill. If thymine is present, GO2, urea, and acetol (CII, COCII, Oil) are formed, and the acetol distills over. Make strongly alkaline with NaOll. Add several drops of o-aminobenzaldehyde ' Evaporate over a flame to half the volume. Gool. Make distinctly acid with liCl and then alkaline with NaliCO, Fliter. A blue fluorescence of 3-oxyquinaldine indicates acetol and the original presence of thymine

(g) ULTRAVIOLET ABSORPTION SPECTRA The availability of convenient and relatively inexpensive ultraviolet spectrometers has made it possible to make most of the quantitative and qualitative measurements of the individual purines and pyrimidines through their strong and characteristic ultraviolet absorption spectra. See also n. 283.

9 Methods for the Separation of Purmes and Parimidines. Procedures which involve fractional crystallization for the separation of purines and pyrimidines have been largely superseded by the techniques of ton-exchange separation or paper chromatography.

(a) CHYSTALLIZATION OF DERIVATIVES OF GUANINE AND ADENINE FROM NUCLEIC ACID Place 5 g of yeast nucleic acid in a 200-mi Erlenmeyer flask, add 100 ml of boiling-hot 5 per cent sulfuric acid and heat carefully over a small flame to bring the nucleic acid into solution. During this heating, which lasts only a minute, the contents of the flask should be kept constantly in motion to avold charring the undissoived nucleic acid. Close the flask with a cork bored with one hole into which is inserted a condensing tube. Immerse in a holling water bath and heat for about an hour. Disconnect the flask and while Its contents are still hot add concentrated ammonia a drop at a time until the fluid is slightly alkaline, and then add 5 ml of concentrated ammonia in excess Guanine is thus precipitated in granular form, while all of the other products including phosphoric acid and adenine remain in the ammoniacal solution.

After several hours, collect the precipitated guanine and wash it with 1 per cent ammonia. Dissolve in as small an amount of 20 per cent sulfuric acid as possible, add a little charcoal, and boil Filter, heat to boiling, and precipitate with an excess of ammonia Filter, dry the precipitate at 40°C, and dissolve it in about 20 parts of boiling 5 per cent hydrochloric acid. As the solution cools, guanine hydrochloride separates as needle-shaped crystals.

⁴ Mix 3 g. of crystalline o-mitrobenzaldeh; de with 50 g. of crystalline ferrous sulfate. Add 75 ml of concentrated ammenta. Heat on a steam bath for one hour. Distill off the c-aminobears! lehyde with steam. The mixture before distillation will keep for two weeks

Combine the ammonlacal flitrates obtained in the isolation and purification of quanine. Filter if necessary. The ammonia is then boiled off and an excess of picric acid added. The reliow precipitate of adenine picrate is collected and dried. The adenine picrate can be recrystallized from 25 per cent acetic acid. Adenine can be recovered from the picrate by acidification with HCl and long extraction with ether, or by the use of an ion-exchange resin which will take up the picric acid.

(b) SEPARATION OF PURINES BY ION-EXCHANGE CHROMATOGRAPHY. Place a loose pad of class wooi in the hottom of a 50 mi. hurette. Add about 25 ml. of water and then sufficient suspension of Dower 50 (hydrogen ion form) to make a resin hed 2 cm, high. Wash the resin into place by filling the huret to the top with water and allowing it to drain. Use a long rod to push another loose vad of class wool down to the top of the resin bed. The rate of flow should be about 7 or 8 ml. per minute when the huret is full, and 2 or 3 ml. per minute for the last few ml. in the buret. The stopcock should always be closed just before the liquid level reaches the top of the resin bed. Arrange a series of eight numhered tubes for collection of fractions from the column.

Charge the column by pouring in a solution of 0.5 mg, of hypoxanthine and 0.5 mg, of adenine in 50 mi, of 0.1 N HCl and allowing it to drain. Wash the column with a few mi. of water and combine the effluents as Fraction 1. Flute the purines from the column with five consecutive 25-ml, portions of 2 N HCl. Then elute with two 25-ml. portions of 4 N HCl for Fractions 7 and 8.

Test for purloes in the cluates by reduction, diazotization of the products of reduction, and coupling with the Bratton-Marshall reagent, as follows, Place about 5 ml. of each fraction in clean test tubes. To equalize the acid concentrations add 2 ml of 6 N HCi to Fraction 1, and dllute Fractions 7 and 8 with equal volumes of water. As controls use 0.025 mg, of hypoxanthine and 0.025 mg, of adenine, each in 2 N HCl, and a 2 N HCl blank, 1) To each tuhe add a plach of Zn dust the size of a match head and place immediately in a holling water bath for 5 to 7 minutes. 2) Cool the tunes by transferring to a heaker of tap water and filter each. 3) To each filtrate add 1 ml. of 0.3 per cent sodium nitrate solution. 4) After 10 minutes add 1.5 ml. of 0.5 per cent ammonium suifamate solution. 5) Add to each 1 ml. of 0 2 per cent N-(1naphthyi)ethylenediamine solution. The hulk of the hypoxanthine (which feads to a pink-purple color) should he in Fractions 2 and 3. A fittle adenine (which yields an orange-pink) will appear in Fraction 6, but most of it in Fraction 7. The pattern of elution from the column depends to a considerable extent upon the length and compactness of the resin bed.

With larger columns operated more slowly, it is possible to effect more complete separation of complex mixtures of purine and pyrimidine derivatives and of isomeric nucleotides, than by the foregoing method. For a discussion of ion-exchange resins and column chromatography, see Chapter 1.

(c) PAPER CHROMATOGRAPHIC SEPARATIONS OF PURINE AND PYRIMIDINE DERIVATIVES Preparation of the paper: On a sheet of filter paper 17 by 26 cm. (Schleicher and Schuli No. 597) draw a light pencil line 2 cm. from one of the long edges. Place single drops of the compounds to be studied at intervals of 2 or 3 cm.

should be prepared fresh each week.

Dowex 50 (H+) 200-100 mesh is prepared by suspension in 5 volumes of water and removal, by decantation, of any very fine material which settles slowly. The resin is then washed by heating 1 hour in 2 N NaOH on a steam bath, then with 4 or 5 changes of 4N HCl (or in a column until free of material absorbing in the ultraviolet) It is then washed by repeated changes of water until the effluent is nearly neutral. Store under 2 volumes of distilled water When well shaken, 6 ml of the suspension will furnish 2 ml of resin

* This solution is stored in a dark bottle, and all solutions for the Bratton-Marshall test

along this line and allow them to dry thoroughly (or dry under an infrared lamp), if more than one drop proves to he needed, the first must be thoroughly dry before the second is added. Each spot should contain about 25 at Roll the paper and staple it to form a cylinder 17 cm high (do not overlap the edges of the paper but allow the staples to serve as links between the edges).

Stand the cylinder of paper in a 10 cm. petri dish filled to a depth of about 5 mm. with the appropriate solvent Cover with a bell or battery jar." When the solvent has migrated nearly to the top edge of the paper, flatten the paper and dry, in a well-ventilated oven if possible.

Inspect the papers under a low-intensity ultraviolet lamps and outline in pencil those dark areas where the compounds absorb or quench the fluorescence of the paper. The distance the compound moves, expressed as a fraction of the distance between the starting point and the solvent front, is known as the Ry value of the compound. For a discussion of paper chromatography, see Chapter f.

1) Spot adenine, guanine, hyporanthine, and mixtures of adenine and guanine and of adenine and hypoxanthine. Develop with n-hutanol saturated with water, with a small container containing a few ml. of conc, NH,OH also placed under the jar. Allow about 3 to 4 hours for development.

2) Spot the same compounds and develop 3 bours in 70 per cent ethanol. This solvent is rapid, but the spots are less discrete. Adenine and hypoxanthine can be differentiated from quanfne, but not from each other.

3) Spot guanine, adenine, cytosine, uracil, thymine, and mixtures of guanine and uracil and of adenine and thymine Develop in isopropanol-2 N 11Cl, for about fe hours. This solvent moves slowly but does permit differentiation of nil these five bases

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f Preferably with a tightly fitting glass plate as a base

Mineralight, 1-41, Ultraviolet I roducts Los Angeles,

This solvent must be prepared fresh each day To 16 4 ml of cone. HCl and 65 ml, of is propanol add water to 100 ml

8

Milk

The first food naturally taken by the newborn baby or the young of any mammalian species is colostrum This is the secretion of the mammary glauds during the earliest phases of factation. Although it may appear before parturition, it is characteristic of the second or third day post partum of the human mother, and is secreted in a total amount of about 100 ml This early secretion is more viscous, richer in protein and salts. and poorer in sugar and fat than milk. The color of human colostrum is lemou yellow while that of cattle is reddish yellow Bovine colostrum, examined just after calving, usually contains vitamin B. in concentrations significantly higher than the 7 µg per liter average found in mature milk 1 Colostrum, unlike mature milk, is heat-coagulable, since it contains high concentrations of globulus sometimes exceeding 10 per cent in cattle Among these globulins have been identified many immune globulins

It has been recognized for more than half a century that newborn mammals acquire immunity to certain infectious by ingestion of colostrum This mechanism appears to be more important in calves than in babies Immune globulius are absent from the blood of newborn calves, but appear three hours after colostrum is fed. This ability to absorb significant amounts of intact protein appears in the newborn of several species, but is limited to their first 24 to 48 hours. There is evidence? that the absorption of immune globulins is effected by the intestinal lymphatics rather than by the portal system The presence of trypsin inhibitor in colostrum's may aid this temporary process of passive alimentary immunization by delaying the digestive bydrolysis of protein

In cattle, precolosirum may be expressed from the udder at about halfterm The globulm content is high, but the normal constituents of milk are deficient. The presence of precolostrum before lactation starts and its dilution with milk as lactation begins explain the composition of colostrum and the short duration of its production at the start of lactation An antibody (agglutinin for Brucella abortus) has been fractionated from the globulias of precolostrum 4

Mature Milk. In human lactation, the secretion of colostrum does not continue beyond the fifth day after childhirth There is a transitional

 [\]unknown Couch Rupel Henderson and Brown J Davry Sc. 34, 749 (1951)
 Combno Roberts and Titchen \underson \unknown \underson \under

period of about five more days during which the components gradually change their proportions to those of mature milk. The fat content of human milk increases during the transitional period and continues to rise for the first two weeks, with no consistent variation after the third week. Increase in lactose content continues for several weeks after the establishment of mature milk flow.

COMPOSITION OF MATURE MILK Representative values in g per 100 ml whole milk*

Constituent	Human	Cow	Goal
Total solids	12 4	12 8	13 6
Protein	1 2	3 3	3 4
Lactore	7 0	4 8	4 7
Fat	3.8	3 8	4 1
Ash	0 21	0 71	0 77

*Summarized from Macy Kelley and Sloan The Composition of Make (Bull Vot Research Council \$113) Washington D.C. 1203

In the dairy cow the change from colostrum to mature milk is a continuous process taking from 5 to 12 days The composition of mature milk is subject to considerable variation, depending upon the breed and the state of nutrition of the dairy animal, the duration of lactation, and the climate Therefore, the accompanying table and similar tabulations purporting to represent the composition of milk must be recognized as simplified collections of averages, deviations from which are to be expected in the analysis of any particular sample of milk. The extent of some of these deviations will be indicated in following sections. For much more detailed tables, including many constituents not considered here, consult the monograph of Macy, Kelley and Sloan, from which the fig ures of the table above were summarized The customary pooling of mar-Let milk tends to minimize variations in composition. In the process of pooling the fat content is usually standardized Since milk is an important article of commerce and since its nutritive value is of importance to the public health standards of composition have been enacted into law Lagal definitions of milk vary in different jurisdictions, but many are consistent with the following

the milks of other species may not be sold simply as "milk" Furthermore, in many jurisdictions, milk of other species is not subject to the requirements of quality and cleanliness imposed by law

Milk Sugar. The only earhohydrate present in nutritionally significant amount in milk is lactose, a disaechande consisting of one glucose unit and one galactose unit with the glycoside link between the earhon 4 of glucose and the earhon 1 of galactose. This glycoside link possesses the beta configuration, so that the full chemical name of lactose is p-glucopyranose—1-(β-p-galactopyranoside). Lactose is a reducing sugar and exhibits mutarotation, since eithou 1 of the glucose portion is free. Like other reducing disaechandes, lactose reduces and Cu⁺⁺ solutions more slowly than do monosaccharides, and hence can be distinguished from them by the use of Barfoed's reagent Lactose forms a specific phenylosazoue, yields a positive muce and test, and is not fermented by the

Lactose is formed by the mammary gland from glucose or glycogen ⁵. The lactose content of milk is not notably altered by changes in the maternal diet or level of blood sugar. Lactose can frequently he detected in specimens of human urine collected post partum or during lactation.

Lactose is converted to lactic acid in the ordinary souring of inilk which is brought about by enzymes of Strepleoceus lactis and numerous other microorganisms. The first step in this conversion is the hydrolysis of lactose by lactase in the surface of the bacterial cell $^{\delta}$ The glucose is converted to glucose-6-phosphate by the usual steps. Galactose is phosphorylated by ATP to α -galactose-1-phosphate in the presence of galactose-hanase. In the presence of undine diphosphoglucose α galactose-1-phosphate is converted to α glucose-1-phosphate, which is converted to glucose-0-phosphate by steps which together with subsequent glycolytic steps, are identical in lactic acid hacteria with those in muscle (see Chapter 10) 7

The percentage of lactose varies significantly in the milks of different species. Human milk has a higher lactose content than that of dury amiliar. The mean value in statistically adequate studies is close to 7 g per 100 ml human milk, with standard deviations less than 0.5 g. The mean for cattle and goats is slightly less than 5 g. lactose per 100 ml milk, with somewhat larger variations. When babies are reared on dury milk, it is customary to add carbohydrate in the form of lactose, sucrose, glucose, or partially hydrolyzed starch (maltose and dextinis).

Lactose is present in no other foodstuff than milk, except of course foodstuffs to which milk or lactose from milk may have been added Compared with the other common sugars, lactose is less soluble and less sweet. The lack of sweetness is considered to be advantageous in the feeding of infants and invalids, since there is less tendency to cloy the appetite Nutritional-Lalance studies unth a small group of children have nudicated

ordinary commercial strums of yeast

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Reithel Fersonal communication
Cori C Personal communication

Mills Breiter Kempster Mckey Pickens and Outhouse J Valention 20 467 (1940)

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that calcium retention is greater on a dict containing 36 g of lactose per

day than on one otherwise equivalent but with no lactose

Milk Lipides Fats occur in milk in the form of suspended globules, of a size easily observed with the low-power microscope. The size of the fat globules in a given specimen of milk is somewhat variable, the largest globules being about seven times the diameter of the smallest Some varia tions in average size of fat globules occur in the different breeds of dairy cattle It is apparent from the large size of the globules of fat in milk that such a suspension of fat in an aqueous system would not be indefinitely stable

I large proportion of the fat of cow's milk rises spontaneously to the top of the container upon standing, and may be skimmed off as cream The separation of cream is sharper and more rapid if the milk is first diluted. The centrifugal separator effects a prompt separation of cream without dilution. A sharp cream line is seldom observed when human or goat's milk is allowed to stand, but the cream can be separated by dilu tion and standing or by centrifugation. This difference in cream-rising, in part the result of larger fat globules in cow's milk, can be altered by homogenization, which is the mechanical reduction of the size of fat clobules

The mean values for fat in human, cow's, and goat's milk (see p 220) are not far apart. In contrast to this consistency of averago values in the three species, individuals vary widely in the fat content of milk, and among cattle and goats there are distinct differences according to breed The milk of Holstein cattle tends to run below the mean value of 3 8 per cent, whereas good specimens of Jersey cows almost consistently give milk exceeding 5 per cent in fat content. The records for fat production, however, are, and long have been, held by Holsteins, since they make up for lower fat percentage by higher volume production of milk Low fat values around 1 per cent and high values around 9 per cent have been occasionally recorded on authenlie specimens of human, cow's, and goat's milk even when proper precautions bave been taken to avoid sampling

The fat content of cream varies according to the method of separation and is usually less than 35 per cent. Butter is a still more concentrated form of nulk fat, conlaining more than 80 per cent. The Babcock test for fat in milk was introduced in 1890 as a practical method for routine use in daines and creamenes. High analytical accuracy was not claimed. In practice, the Babcock test has been found to give results only slightly (00) to 007 per cent) higher than the standard Roese-Gottlieb etherextraction incthod, or its mechanized equivalent the Mojonnier method The modification of the Babcock test described in the experimental sec tion of this chapter is not official and would not be acceptable in estabh hing butterfat records or in court actions. In such situations the analyst should familiarize himself with the regulations governing butterfat analyes in his particular state or territory

The inglycendes which compose milk fat contain a complex mixture of fatty acids with olcie and palmitic acids quantitatively predominant The fat of human milk is characterized by a somewhat higher proportion of oleic acid, and a relative deficiency of short-chain fitty acids (butyric, caproic, caprylic, and capric) as compared to goat's or cow's milk. Comparing the latter two species, cow's-milk fat has much more stearic acid, somewhat more butyric acid, and notably less caproic, caprylic, and capric acids than goat's milk. These differences in fatty acid distribution are probably of little nutritional significance, but become apparent when one compares the hutyric odor of rancid cow britter with its goat counterpart. Of greater significance are certain fatty acids with more than one double hond. Rats develop dermatits when fed an otherwise adequate diet deficient in such multiply unsaturated fatty acids. Hansen and Burr's list inolcie acid, linolenc acid, and arachidonic acid as normal dietary components which in very small amounts will relieve this deficiency. The milks of all three species discussed contain arachidonic acid, and the concentration in goat's and cow's milk is somewhat higher than in human milk. The presence of linoleic acid in milk has heen verified.

Cholesterol is present in small amounts in milk. The mean value for con's milk is 11 mg per 100 ml. Both fatty acids and cholesterol are synthesized in the lactating mammary gland, as demonstrated by the rapid incorporation into these milk lipides of injected carboxy-C¹⁴ acetate, both in the intact lactating goat¹² and in the perfused isolated udder of

the cow 13

Milk Proteins. Human milk is notably low in protein coutent, with a nean concentration of 1.2 g per 100 ml and rare variations helow 1 or hove 2 g. The mean values for cow's and goat's milk are 3.3 and 3.4 g per 100 ml, most analyses falling within the limits of 2 and 5 g. In the preparation of formulas for feeding young infants dilution of darry milk is a customary but not universal practice. Although such dilution, together with the addition of carbohydrate, yields a mixture more closely resembling liumau milk, the emphasis in present day infant feeding is upon meeting the nutritional needs of the infant rather than upon close imitation of the composition of human milk.

The chief and characteristic protein of milk is casen, a complex of phosphoproteins which makes up a third of the protein of human milk three-fourths of the protein of goat's milk, and a little over five-sixths of the protein of cow's milk. Casem is definitely not a single molecular species, since three electrophoretic peaks have been distinguished (see p. 7). Fresh milk, at its pH of 66 df 0.2, has as its chef colloidal component a complex of calcium casemate and phosphate. If the milk is hrought by natural souring or by cautious acidification to pH 4.55, the isoelectric point of casem, this complex is broken up and casem is precipitated. The product of such a precipitation may be designated isoelectric casem. Casem can also be precipitated by the action of certain proteolytic curymes, the best known of which is remain (remet to choese makers)

Hansen and Burr J Am. Med 4ssoc 132, 850 (1945)
 White and Brown J 4m Oil Chemists Soc 26 335 (1949)
 Natal Mickelsen heys and Petersen J Vulriton 36 490 (1948)

Popjak, French and I olfey, Biochem J., 48, 411 (1941)
 Cowie Duncombe Folley, Glascock Massart Feeters and Popjak Biochem, J
 Glu (1951)
 Lovico and Haller J. Phys. & Colland Chem. 51, 409 (1947)

obtained from the fourth stomach (abomasum) of nursing calves Rennin does not occur in the human being but other enzymes, in particular pepsin and thymotrypsin, carry on the same function Crystallized rennin has been shown to be a proteolytic enzyme which, in causing milk to curdle, hydrolyzes one peptide bond per 10,000 in casein 15 The precipitate resulting from the coagulation of milk by proteolytic enzymes is therefore not identical with isoelectric cascin, but has been subjected to a mild degree of proteolysis This product is called paracasein, and at pli values above the isoelectric point, is precipitated as calcium paracaseinale Further proteolysis of casein yields the usual intermediate products of protein hydrolysis, except that phosphopeptones containing phosphoserine have been isolated from tryptic digests of casein. The caseins of milks of different species are not identical in composition

Except for some cottage cheese (Dutch eheese or schmierkase), which is usually crude isoelectric casein, the varieties of cheese on the American market are made from the calcium paracasemate obtained by rennin coagulation of milk Cheddar is obtained by rennin eoagulation of milk preferably pasteurized while fresh and then slightly soured by a starter of lactic acid hacteria Later processes include cutting the curd, heating to about 40° C, straining, and cheddaring, which consists of matting the theese together by piling, cutting, and repiling During the final curing process, Lactobacillus cases overgrows other organisms and produces th characteristic flavor Swiss cheese is started in a similar manner, but 15 cultured with Propionibaclerium shermanii to produce the "eyes" and the sweetness There is no cheddaring process, but the curd is cut and cooked firm Curing takes three to six months, during which time hacteriologi cal errors may occasionally produce such anomalies as "blind" cheeses, "stinkers," and "blowers" Although in France Requefort is often made from cwc's milk, American blue cheeses are made from cow's milk, and are started like cheddar. The cut curd is drained, placed in small perforated forms, and sprinkled with Penicillium requeforti Further ripen ing and proper mold-mottling requires earefully controlled curing for six to nine months The ripening of Limburger depends upon Bacterium linens and other naturally occurring bacteria of the milk acting upon a rennin curd produced from fresh milk without a starter, and takes about two months Yeasts growing on the cheese surface provide essential nutrients for B linens 16 Camembert is the product of natural hacteria working inside the cheese while the mold Penicillium camemberti grows on the outside

Casein originates in the lactating mammary gland. Certain of its component amino acids have been shown by isotopic identification to be taken up from the free amino acids of the blood, 17 and not from plasma proteins The pho-phorus of cascin has similarly been shown to be derived from the morganic phosphate of the blood. In the goat, about two bours are required for the transport of these substances from circulating blood to the casein of the milk

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Nitschman and Varin Hels chim. acta 24, 1421 (1951)
 Purko Nelson and W. od. J. Dairy Sci. 14 1139 (1151)
 Barry J. Biol. Chem. 195, "95 (1242)

Chap 8 Proteins other than casein constitute more than half of the proteins of human milk, and an appreciable fraction of the proteins of dairy milk True globulns have already been mentioned in connection with colostrum After the removal of easein from cow's milk by isoelectric precipitation, the remaining whey is found to contain 0 6 to 0 7 per cent protein, of which roughly one-fourth is globulin, as shown by its being precipitated when the whey is saturated with magnesium sulfate at pH 7, one-half is albumin, which is not so precipitated but is heat coagulable, and the remaining fourth is proteose, which is not heat-coagulable From the "albumin" fraction has been crystallized a β-lactoglobulin which is probably identical with other preparations described as lactalbumins. The amino acid composition of β -lactoglobulin is quite different from that of any plasma protein, but the immune globulins of colostrum are similar in composition to their plasma counterparts

Among the proteins of milk are included numerous enzymes The pattern of enzymes is quite different in milks of different species Aldolase and vanthine o'udase (Schardinger enzyme) are present in cow's milk but not in human milk Catalase and peroxidase are present in raw milks Human milk, particularly rich in amylase, promptly liquefies stareb paste The lipase of human or bovine milk brings about a slow liberation of free fatty acids from milk fat Cow's milk collected in mid lactation contains from 80 to 120 King Armstrong units of alkaline phosphatase (phos phomonoesterase) per 100 ml, compared with a range of 0 7 to 16 2 units in human milk and a mean of 48 units (Kon and Mawson) The phosphatase content of cow's milk has been made the basis of tests for adequate pasteurization, since the enzyme is readily destroyed by heating

ESSENTIAL AMINO ACIDS IN WHOLE MILK PROTEINS (Expressed as per cent of total protein (\ = 16))* zlh

ATT and ag ner	cent of total protess	
(Expressed as per	Human Wilk Proteins	Cows Mill Proleins
Arginine	4 3 2 8	4 0 2 8
Histidine Isoleucine	7 2	7 1 10 4
Leucine	9 8 7 2	83
L) sine	2 2	3 0
Methionine Phenylalanine	5 6	5 2 4 5
Threonine	46 19	1 4
Tryptophan		6.7
Value	lerived largely from Illo	ck and Bolling

These representative data are derived largely from Block and Bolling The Amino Acid Ameso representative data are derived largety from more and county, we amino acid Composition of Proteins of d Foods Charles C Thomas Springfield III 1951 Te N = 16. convention is used in detary calculations although for milk proteins (large), casein) N = 13.7 (i.e. protein = N × 6.38) is more accurate For additional data on individual multimilk proteins see table on p 122

The biological value of a protein food is defined as the ratio (expressed as percentage) of the food mtrogen retained to that absorbed (see p 1047) It is usually determined by feeding experiments on growing rats A high value indicates adequacy in content and availability of so-called essential amino acids. The biological values of human and cow's milk are identical within limits of experimental error, the numerical range being \$5-90. This high biological value is confirmed by direct analysis of milk proteins for escential amino acids. The essential amino acid milks protein based on the relation hip between its essential amino acid content and that of whole-egg protein (ascumed to equal 100) is 90, the corresponding in dex for human milk protein is 93. These index values are closely correlated with the biological values. ¹³ The table on p. 225 demonstrates the adequacy of whole milk in those amino acids generally regarded as indispensable in the human diet. This question is more fully discussed in Chapter 33.

Inorganic Vilk Components. The crude figures for ash in the table on p 220 indicate that the concentration of total minerals is more than three times as great in the milk of dairy animals as in human milk The important bone-mineral elements, calcium and pho-phorus, are both preent in milk in nutritionally significant amounts and in a ratio suitable for

effective absorption and utilization

Calcium is present in human milk in concentrations which vary through a range of 10 mg per 100 ml on either side of a mean of 30 mg. Cow's or goat's milk averages close to 130 mg. per 100 ml. The calcium of milk 13 precent in part as calcium ion, and in part bound as a colloidal complex with protein and phosphate

Phosphorus has a mean value of 13 mg per 100 ml in human milk, with a standard deviation of 19 mg (Kon and Mawson), but is found in amounts averaging 100 mg per 100 ml of dairy milk. 70 to 80 per cent of the total pho phorus being in the form of inorganic phosphate. The remainder includes phosphoprotein, phospholipide, and ester phosphorus A variable portion of the inorganic phosphate is separable by ultrafiltration as a part of the colloidal calcium caseniate-phosphate complex.

During that period of early life when milk is the only food taken, potassium is equal in importance to the bone minerals At all ages, the body contains more potassium than sodium, but the difference is greater with growth Human milk contains approximately 11 milliequivalents of potassium and a milliequivalents of sodium per liter this ratio seems to be optimal for the growth of the mfant Dairy milk contains these ions in comparable ratio, but in concentrations about three times higher Chloride ion is present in highly variable amounts. Magnesium averages 4 mg per 100 ml of human milk, and three to four times this concentration in dairy milk. The sulfur of milk is almost entirely in the form of the sulfur containing amino acids of milk proteins and averages 30 mg per 100 ml of cow's milk, and about half this value in human or goat's milk.

In addition to those mentioned above, a number of other elements are present in small and variable amounts. Vilk is notoriously deficient in iron. Inalytical figures from different laborationes are usedly divergent, but for cow's milk a concentration of 0.32 mg. per kg. is representative. 19 The iron content of milks of other species is comparable. The full term numan infant born of an adequately nourshed mother has a store of iron.

¹⁶ Over J Am. Dudd Assoc 27 3x6 (13x1)
¹⁹ Johnston Food Les ? 212 (1344) Johnston Gellman, and Strom J Bud. Chem-173, 343 (1945)

for about six months of growth on a milk duet without developing an iron-deficiency anemia After six months, supplementary iron-containing foods should be introduced into the diet Copper occurs in raw cow's milk in about the same concentration as iron Higher concentrations may result from contact with copper or brass during pastcurization or other processing Copper in excess of 05 mg per kg produces a tallowy flavor and loss of ascorbic acid in milk Average values for other trace elements in cow's milk are given by Archibald20 as follows

Vianganese	22	μB	per	liter
Zinc	3900	46		**
Cobalt	0 6	"	46	16
Molybdenum	73	**	44	44
Nickel	none			

The presence of jodide has been demonstrated in the milk of women and of animals Electrophoretic evidence has indicated that milk contains an iodinated protein 204

Vitamins in Milk, Milk and milk products are outstanding sources of

certain vitamins, and less reliable for the supply of others

Thramine is present in human milk in amounts which roughly reflect the vitamin B1 intake of the mother At moderate levels of intake, about 10 per cent of ingested thiamine is secreted into the milk (Kon and Mayson) The concentration of total thiamine (in the milk of mothers whose intake is between 15 and 2 mg) usually lies between 14 and 19 µg per 100 ml In early lactation, from 50 to 90 per cent of the thiamine of buman milk is in a combined form which is not reactive in the thiochrome test, but becomes reactive after enzymic hydrolysis with takadiastase, which contains numerous enzymes including phosphatases and proteases In general, high content of combined thiamine is associated with low values of milk phosphatase Both free and combined thiamiue are measurable by bioassay. The proportion of free thiamine increases with the duration of buman lactation

The milk of goats and cattle contains somewhat more than twice the concentration of thismine in human milk. About half is in the form of free thiamine in cow's milk, and about one sixth in goat's milk. Thiamine pyrophosphate added to cow's milk is promptly hydrolyzed in the presence of the enzymes pyrophosphatase and phosphatase A portion of the combined thiamine in cow's milk is bound to protein and liberated by hydrolysis with pepsin In the cow, total thiamine of the milk is higher in early lactation than in the later months As in human milk, the thiamine of cow's milk varies with the amount supplied in the diet, but the cow has the additional possibility of supplementation by thiamine formed by microbiological synthesis in the rumen. One quart of milk contains about one-fourth the daily human adult requirement of thiamine

Riboflavin is the one vitamin for which dairy milk and milk products are the richest cveryday source, exceeded in riboflavin concentration only hy liver and yeast. The source of this ahundance of riboflavin is bacterial

Archibald J Dairy Sci., 34, 1026 (1951)
 Middlesworth Tuttle, and Threlield Science 118, 749 (1953)

action in the rumen. In cattle, a day's output of riboflavin in the milk has been found to be up to ten times the dietary intake. Alost samples of darry milk contain between 100 and 200 µg of riboflavin per 100 ml Cheese and other inilk products, including evaporated milk, are excellent sources of riboflavin. If milk and milk products are excluded from the usual American diet the riboflavin requirement is not met. The riboflavin of milk and milk products is as completely available to the luman organism as is riboflavin taken in simple water solution. This is not true of the riboflavin in some other common foodstuffs.

The riboflavin content of human milk is much lower than that of cow's milk, usually between 20 and 30 gg per 100 ml if no supplementary riboflavin is ingested. Since the contribution of riboflavin to the human organism from bacterial synthesis in the large intestine is minimal, the riboflavin content of human milk is determined by the riboflavin of the dict.

Nicotinic acid is present in cow's milk in a concentration usually slightly less than 100 µg per 100 ml. Somewhat higher concentrations are reported for goat and human milk. It is obvious that the recommended daily adult intake of 10 to 15 mg could not be met with milk alone, and one might wonder how the nursing young avoid deficiency. A part of the answer is that many animals, including man, can meet part of the nicotinic acid requirement by synthesizing it from tryptophan. Pellagra-producing dicts have been characterized by low content of tryptophanias well as of meotinic acid. Milk, although low in mechanic acid, is high in tryptophan (40 to 50 mg per 100 ml in cow's milk)

Cattle appear to be able to meet their entire requirement for panilothenic acid by bacterial synthesis in the rumen Dairy milk contains 300 to 400 μg of pantothenic acid per 100 ml, human milk about half this amount. The pyridacial group of vitamins (vitamin Be) is present in milk in variable amount, most reports giving mean values between 50 and 70 μg per 100 ml for cow's milk Bothn is present in milk in quite variable concentrations up to 10 μg per 100 ml. The mean folic acid and vitamin B₁; contents¹¹ for human milk have been reported as 0.71 and 0.41 μg per liter, for cow's milk, 13 and 6.6 μg per liter and for goat's milk 2.7 and 0.12 μg per liter All the B vitamins mentioned in this paragraph are those of which an uncomplicated detary deficiency has not been observed in man, and which in large part may be supplied to man by bacterial synthesis in the intestine hence the values given are of more academic than nutritional significance

The ascorbic acid content of cow's and goat's milk, when fresh, is about 2 mg per 100 ml, which decreases to less than 1 mg by the time the milk is made available through usual retail outlets ¹² For infants, and for adults living on milk diets supplementation with ascorbic acid is necessary for optimal intakes. Human milk depending somewhat upon the nutrition of the mother usually runs higher in ascorbic acid than the milk of dairy animals. On intakes of vitamin C ranging from 43 to 106 mg

 ¹¹ I verson learson and Matteson J Vutrition 46 45 (1952)
 12 Collins Harper Sci rether and Livel jets J Vutrition 43 313 (1951)
 14 Holmes J in Dictair Assoc 27, 578 (1951)

daily, 10 mothers were found to put out from 19 to 46 mg of assurbed acid in the milk per day, at concentrations ranging from 19 to 58 mg

per 100 ml (Kon and Mauson)

Vitamin Å, heigh fat-soluble in contrast to the vitamins previously mentioned, is found in association with milk fat. Human milk may contain from 50 to somewhat over 200 units of vitamin A per 100 ml, the concentration of the vitamin being correlated with the fat content of the milk. The carolenoids which are responsible for the yellow pigmentation of milk fat are to a varying degree providenies. 4 Human milk contains from 7 to 25 µg of carolenoids per 100 ml. The milk of cattle contains vitamin A and provitamins A in concentrations which, though comparable to those in human milk, are highly variable, depending upon the composition of the feed.

Vitamin D, under optimal conditions, may be present in human milk, dairy milk, or milk fat in amounts adequate to prevent rickets in infinits and children Since this is often not the ease, supplementation with vitamin D is standard practice in those parts of the world where exposure of infants, mothers, or dairy animals to sunlight is indequate Milk is often

enriched with vitamin D to provide 400 units per quart

Vitamins K and vitamins E are present in measurable amounts in liuman and dairy milk. The concentrations of these substances in milk are not significant in human nutrition.

MILK AND HUMAN NUTRITION

The suitability of milk and milk products for human food is a logical consequence of their chemical composition. The history and experience of mankind give further evidence of the value of dairy animals to our well-being. The cow produces a protein food of high biological value and produces it with an extraordinary economy which results from her ability to utilize a high proportion of rough feed in her diet. As an extreme case, she can survive and produce milk on forage alone, although this is not the way to maintain a high milk production. Protein foods are, of course, necessary for man at all ages, but their lack is most disastrous during early childhood shortly after wearing Kwashiorkor, or malignant malnutrition, occurs in children in those areas where the proteins of high biological value supplied by such foods as milk, meat, and egg are lacking, and where starchy foods are the dietary staples 24 This disease is characterized by extreme fatty degeneration of the liver, with permanent scarring (cirrhosis) in those who survive Although kwashiorkor is a disease of multiple dietary deficiencies, usually with adequate caloric intake it responds poorly to vitamin therapy, and is best treated or prevented by protein foods

Milk has a justified reputation as a "protective" food, meaning that the protects the individual from several possible dictary deficiencies, in cluding those of protein, calcium, and inhofiavin Like many other good things, milk can be used unwisely Older children require more protein than is supplied by a quart of milk, and it is wise not to depend upon milk.

²⁴ Meiklejohn and Passmore Ann Rev Med., 2, 129 (1951)

alone as a protein source, but to supplement it with more concentrated protein foods. It should also be recalled that milk is not a dependable source of iron, thiamine, ascorbic acid, or vitamin D (except vitamin D milk) in the diet of growing children. Iron deficiency anemia is quite characteristic of infants who have been kept too long on unsupplemented breast or bottle feeding.

The use of human mill has obvious advantages in the feeding of human infants, including optimal adaptation to nutritional needs and almost complete freedom from bacterial contamination. In lactation, the first week is entical for the mother, who should avoid overeating or other possible causes of digestive disturbances. Later, the duet should be ample and nutritious, but there need be no excessive gain in the mother's weight. The addition of one quart of dairy milk to her usual diet is a common and usually satisfactory recommendation. The average daily production of milk by the human mother ranges from around 300 ml in the first week to 900 ml in the twentieth week. The variation from these mean figures is extreme, and outputs of more than a gallon a day have been reported

Cow's mill as the basis of most prepared foods for infants. The reduction in the mortality of bottle-fed habies has heen one of the untrumpted trumphs of proventive medicine. As recently as 1922, the mortality of bottle-fed habies was four times that of breast-fed habies. At present there is no significant difference. Pasteurization (heating to 143° F for 30 minutes) destroys pathogenic hacteria without coagulating the whey proteins. This is an adequate method, when combined with proper handling after pasteurization, of rendering the milk suitable as a food for adults. It climinates the possibility of transmission of brucelloss or borne tuberculosis to human beings. It is not an adequate treatment of milk for young babies, for whom all milk, whether raw or pasteurized, is customarily boiled. This reduces the toughness of the casein cut (see p. 360) and kills all bacteria. There is no nutritional disadvantage to the boiling of milk other than the complete destruction of ascorbic acid, which is customarily given in a supplementary food.

Goat's mill is used as a food to a limited extent in the United States A considerable amount is produced in small dairies chiefly for home use When goat's milk is available on the market the price is usually higher than that of cow's milk. The nutritive value of goat's milk is so close to that of cow's milk that it is impossible to make a choice on this basis alone Goat's milk has often been used successfully in the feeding of infants or children who show allergy to cow's milk Many other therapeutic claims have been made for goat's milk, but acceptable evidence for them is lacking. The claim of easier digestibility compared to cow's milk has ome basis in fact since the fat particles of goat's milk are smaller, and the curd tension (toughness of precipitated casein) is less as compared with unhomogenized cow's milk. One often mentioned disadvantage of goat's milk is apparently fictitious-so-called "goat's milk anemia" In experiments with both rats and babies no difference has been found in the outcome of feeding with cow's and goat's milk. Either is nutritionally satisfactory and both require supplementation with iron and with vita mins C and D Under the very best stabling conditions for both species, as for example for the experimental herds at Beltsville, it is possible to keep goats cleaner than cows, hence the bacterial counts of goat's milk are lower under such test conditions. Under ordinary stabling conditions, goat's milk has no less opportunity to be contaminated than cow's milk Goats are not susceptible to boxine tuberculosis, but otherwise any milk-horne disease can be carried as effectively by goat's milk as by cow's milk, hence pasteurization or boiling is as necessary for the milk of one species as for that of the other Inspection of cow's milk and of the dairies where it is produced is almost universal in American communities, but goat's milk is often not included in the regulations governing such inspections

Etaporated milk is prepared by concentration under reduced pressure to slightly less than half the original volume. The concentration of dissolved solids is therefore slightly more than twice that of the original milk. For purposes of infant feeding, exaporated milk may be used in half the amount specified for whole milk, reconstituted to the specified volume with holled water. Most brands of evaporated milk sold at retail are fortified to provide 400 U.S. P. units of vitamin D₁ per reconstituted quart, and this is indicated on the label. Evaporated milk is sterilized at high temperatures after being sealed in cans, and under present-day conditions of manufacture is rarely subject to spoilage from incomplete sterilization. Even though the milk is sterile, undesirable alterations of viscosity may occur on storage, sometimes with separation of proteins into an insoluble gel. These changes are unlikely if evaporated milk is used within two years, and if it is stored at temperatures below 60° F. *50.

Canned fresh whole milk which will keep four to six months without refrigeration, is sterilized by hringing its temperature almost to 300° F for a few secouds in a specially designed heat-exchanger. The canning is done under sterile conditious. This type of canned milk has no cooked flavor. At the time of this writing, this product is not sold extensively at retail in the United States, but is processed chiefly for the armed services.

The original condensed mulk (1858) was concentrated by evaporation under reduced pressure and preserved by the addition of sucrose. It was not originally sold in cans, but in bulk after the manner of sale of mulk at that time. For many years this product his heen available at retail, canned and hest-sterilized. The concentration of milk components in present-day condensed mulk is a little higher than in evaporated milk, and the percentage of added sucrose is about 40 per cent.

The better grades of dried mill are prepared by very rapid drying under reduced pressure, which preserves the vitamin potency as well as other nutrients. Such products are satisfactory for human food and for infinits' formulas, when produced under proper sanitary conditions. Che uper grades, intended for animal food, are produced under less strict standards of quality and sanitation.

Storage of fluid milk or of milk concentrates at very low temperatures (-5° to -15° F) is a satisfactory method of preservation which does not produce undesirable changes in flavor or physical properties "The use

Webb Deysher and lotter J Dairy Sci. 34 1111 (1951)
Trac: Hetrick and Krienke J Dairy Sci. 33 832 (1950)

of eatbode rays has been found effective in sterilizing milk, and is claimed to be without effect upon flavor 27

State laws in the United States require that ice cream contain a specified percentage of milk fat, the percentage varying from 8 to 14 in differ ent jurisdictions Frozen preparations with lower fat content are sold as

ice milk or sherbet

Determining the freezing point of milk is a simple physical test which reveals dilution with water Observation in a mixed milk specimen of a freezing point higher than the limiting value of -0 520° C indicates that such dilution has been made. If the sample is from a single cow, there is a remote chance that this limiting value may be exceeded without the addition of water If the sample is a mixture of the milk of six or more cows, a freezing point above the limiting value is definite evidence of watering, and if the sample is from forty cows or more, a freezing point above -0 540° C raises strong suspicion of watering 28

Economics. The milk-producing cows of the United States number close to 25 000,000 and their output is estimated at over 14,000,000,000 gallons per year, which brings to American dairymen an annual gross in come of more than \$3,760,000,000 Estimates released by the Vill In dustry Foundation apportion the 1950 milk output as having been used

as follows

Fluid milk and cream	
sold in cities and villages	37 39
used on farms where produced	10 19
Butter	
creamery made	22 79
farm made	4 34
Cheese	9 5
Ice cream	5 1
Evaporated condensed and dried milk	6 4
Milk fed to calves on farms	2 7
Milk for other uses	1 0

The figures above do not tell the whole story, since skim milk, buttermilk, and whey remain after eream, hutter, and cheese have been made Lactose for use in the manufacture of food and medicinal products is separated from such residues and marketed at a rate of 40,000,000 pounds a year in the United States About half this amount of casein is similarly obtained, and is used for human and animal food, and in pharmaceutical, paper, paint, and adhesive products Hydrolyzates of casein are used in special diets, in food flavoring and as nutrients in industrial microbiology The whey proteins are also separated, and are used in food products and for many industrial purposes

EXPERIMENTS ON MILK

1 Phosphatase Test for Pasteurization (Cornell Method?) Principle The destructive effect of heat on the natural phosphatase in raw milk is used as the basis for testing the efficiency of pasteurization Phosphatase activity is measured by the

n Chem Eng Verz 29 2818 (1921)

Lythgos J Assoc Offic Agr Chemids 55 442 (1952)

Koskowsky J Davy Sci 34, 1151 (1931) Reproduced by permission of the author and the Journal of Dairy Science

hydrolysis of disodium phenylphosphate and colorimetric estimation of the rile ised phenol Dairy product samples are incubated with the substrate buffered at pill 9 1 to 9 7, and then the protein is precipitated. After filtration the filtrate is brought back to an alkaline reaction with carbooate and BQC^D is added. Blue color is produced if sufficient phosphatase was present in the milk to split the disodium phenylphosphate substrate during incubation. The procedure as given here applies to milk. With tumor modifications (given in the original paper) the Cornell test can be applied to other dairy products.

Procedure ³¹ a Samplino and Neubaring (Long Muthop) For milk and other fluid dairy products, 1 ml of milk or milk product is transferred to a 25 × 150 mm test tube Add 10 ml of warm (40° C) carbonate buffer substrate and 4 drops of U S P chloroform A piece of parchment paper is fitted over the tube with the aid of a rubber band, and the milk is incubated at 32° to 37° C for 18 to 24 hours

h Propertation After incubation, 1 mi of trichloroacetic—HCl precipitant is slowly added to the time the resulting protein precipitate is filtered off through Whatman No. 42 paper (11 cm)

c COLOR DEVELOPMENT Five ml of the clear filtrate is pipetted into a 16 × 150 mm test tube One ml of CuSO, "Caigon" solution and 5 ml of 8 per cent Na₂CO₃ are added Then 2 drops of BQC solution are placed in this solution The tubes after mixing are inserted in a water bath at 37° C for 15 minutes Color development is measured after this interval against suitable color standards or in a colorimeter

INTERPRETATION OF RESULTS All final color readings are after consideration of coutrol values multiplied by a factor of 1 2 to convert to µg phenol per 0.5 ml \ny

10 Sec 2 6-Dibromoquinonechlorounine solution (BQC) under footnote 31

31 Reagents and Materials

Carbonale Buffer Substrate Dissolve 11 50 g anhydrous NarCO: 10 15 g anhydrous NarCO: and 1 09 g pure disodium phenylphosphate in water and make up to 1 liter (pH = 9 80)

Trichloroacitic—hydrochloric acid precipitant Dissolve 25 g trichloroacetic acid crystals in water make up to 50 ml with water add 50 ml cane HCl (approx 36 per cent) and mix thoroughly

Sodium carbonate solution (8 per cent) Dissolve 80 g anhydrous Na CO4 in water an i make up to 1 liter

Copper sulfate Calgon soletion Dissolve 500 mg CuSO. 5H2O and 20 g Calgon

(sodium hexametaphosphate) crystals (tech) in water and make up to 1 liter 2-0-Disconvolutionecolorousine soft ton (BOC) Dissolve 50 mg BQC in 10 ml absolute ethyl or methyl alcohol and store in dark bottles.

Color standards Vale the following solutions as preliminary to making standards-

a Stock phenol solution Dissolve 1 g phenol crystals in water and make up to 1 liter

b Buffer solution Make 1 liter of carbonate buffer containing 11 50 g NarCO; and 10 17 g NaHCO;

c. Diluted phenol solution Using 4 ml of stock phenol solution (a) make up to 500 ml with buffer solution (b) This solution contains \$\mu_{\text{pr}}\$ phenol per ml
Preparation of color standards Place in clean 16 X 150 mm test tubes 0.5 to 5 ml por

Preparation of color standards Place in clean 16 × 150 mm test tubes 0.5 to 5 ml por tuns of Diluted Phenol Solution (c) Add enough Buffer (b) so that total volume of inquid in each tube is 10 ml Then add 1 ml of copper sulfate—Calgon solution to each tube Finally add 2 drops to BQC and 4 drops to B P. chloroform and mix Let stand for 15 innuites at 37° C. Seal tubes with paraffin wax and store in refrigerator. Tubes containing 0.5 10 15 2.5 and 50 ml portions of diluted phenol solution will produce standards of 4.0 8.0 120 2.00 and 5.0 mg respectively after final color development. Color standards of 20 and 5.0 mg can be easily obtained by mixing 4.0 and 2.0 mg solutions with out color development and diluting these with sufficient Buffer Solution (b). Color is then develope 1 m 10-ml portions by adding copper sulfate—Calgon solution and BQC

To obtain alcohol color standards o mi of but; i alcol of are added to a duplicate series of final squeous color standards. The tubes are inverted ten times to extract color and the

cork stoppers are sealed with war.

value over 5µg per 0.5 ml milk or other fluid dairy product for the long procedure, or 2 µg per 0.5 ml for the short metl od is tentatively considered to indicate either underpasteurization or the presence of raw milk products or a combination thereof the short method even to the extent of using samples of the same size, is conducted in the same way as the long method except that (a) incubation is carried out for 1 hour at 37° to 38° C, and (b) chloroform is omitted

CONTROLS FOR LONG AND SHORT METHODS One mi of units, preferably from pasteurized stock is placed in a tube heate 1 to 78° C (170° F) for 15 seconds in a water bath and then cooled immediately. This heated control is tested by the same method

as employed for samples of unknown history

ALCOHOL EXTRACTION If necessary butyl aleol of extraction may be used in either the long or short method especially on critical values. Five ml of n butyl aleolol is added to the test tube containing the aqueous colored solutions, and the latter is inverted ten times. The clear layer appears without centrifuging and is compared against alcohol standards. Alcohol extraction is preferred when a colorina ter is not used as aqueous phenol standards for all incthods deteriorate as a rule relatively randily.

- 2 Influence of Gastric Rennin on Milk Prepare a series of five tubes as
- foliows

 a 5 mi of fresh muk + 0 2 per cent HCI (add drop by drop until a precipitate forms)
 - b 5 ml of fresh milk + 5 drops of rennin solution 25
 - c 5 ml of fresh mlik + 10 drops of 0 5 per cent Na; CO;
 - d 5 ml of fresh mlik + i0 drops of ammonium oralate solution.
 - e 5 mi of fresh milk + 5 drops of 0 2 per cent liCi

To each of the tubes (c), (d), and (e) add 5 drops of rennin solution Place the whole series of five tubes at 40° C and after 10 to 15 minutes note what is occurring in the different tubes. Give a reason for each particular result.

3 Preparation of Casein # Into a 600 ml beaker introduce 200 ml of skimmed (or centrifuged) milk Add an equal volume of water Add from a pipet very carefully, drop by drop with thorough stirring, 10 per cent ilCl until a flocculent precipitate forms (Casem precipitates best at a point slightly more acid than its isoelectric point of pli 455) From 3 to 5 ml of acid are commonly required In milk, casein functions as an acid and exists as K and Ca caseinates, from which compounds it is released by the acid As the isoelectric point is passed, however, the casein begins to function as a base and go into solution as casein hydrochioride fience excess of acid must be avoided if too much acid be added, run in, drop by drop, 10 per cent NaOil solution until precipitation occurs and a clear supernatant fluid is obtained Allow the precipitate to settle decant the supernatant fluid, and reserve it for use in later (4-6) experiments Filter off the precipitate of caseln and remove the excess of moisture by pressing it between filter papers Transfer the caseln to a small beaker, add enough 95 per cent alcohol to cover it, and stir for a few moments Filter, and press the precipitate between filter papers to remove the aicohol Repeat the extraction with aicohol, making sure that the casein is in a finely divided condition. Transfer the casein again to a small dry beaker, cover the precipitate with ether and heat on a water bath, with any flames turned out, for ten minutes, stirring continuously. Filter (reserve the filtrate), and press the precipitate as dry as possible between filter papers. Open the papers and allow the ether to evaporate spontaneously. Grind the precipitate to a powder in a mortar. Upon the casein prepared in this way make the following tests:

- a. Somether. Try the solubility in water, sodium chloride, dilute acid, and alkall.
- b. Millov's Reaction. Make the test according to the directions given on p. 169.
- c. CYSTINE AND CASTEINE SULFUR Test for cystine and cysteine sulfur according to the directions given on p. 168.
- d. Fusion Test for Phosphorus Test for phosphorus by fusion according to the directions given on p. 212.
- 4. Coagulable Proteins of Milk. Place the filtrate from the original caseln precipitate in a casserole and heat on a wire gauze over a free flame. As the



Fig. 63, LACTUSE

solution concentrates, a coagulum consisting of lactaibumin and lactoglobulin will form. Continue to concentrate the solution until the volume is about one-half that of the original solution. Filter off the coagulable proteins (reserve the filtrate) and test them as follows:

- a. Millon's Reaction. Make the test according to the directions given on p.
- b. CYSTENE AND CYSTEINE SULFUR. Make the test according to the directions given on p. 168. Do the coagulable proteins differ from caseln in their reaction to this test? Why?
- 5. Detection of Calcium Phosphate. Evaporate the filtrate from the coagulable proteins, on a water bath, until crystals begin to form. It may be necessary to concentrate to 15 ml. before any crystallization will be observed. Cool the solution, filter off the crystals (reserve the filtrate), and test them as follows:
- a. Microscopical Examination. Examine the crystals and compare them with those in Fig. 64.

- h Dissolve the crystals in nitric acid. Test part of the acid solution for phosphates. Render the remainder of the solution slightly alkaline with ammonia, then acidify with acetic acid and add ammonium oxalate. Framine the crystals under the microscope and compare them with those in Fig. 230, p. 855
- 6. Detection of Lactore. Concentrate the filtrate from the calcium phosphate until it is of a syruplike consistency, and pour it into several volumes of acctone to precipitate the lactose in crystalline form
- a Micro-on real Tameation Examine the crystals and compare them with those in Fig. 63
 - b Bistour That Try Benedict's test upon the mother liquor.
- c Phenythyphazine Test Apply the phenylhydrazine test to some of the mother liquor according to the directions given on p. 63.
- 7. Milk Fat Evaporate the ether filtrate from the casein (Fsp. 3) and observe the fatty residue. The milk fat was carried down with the precipitate of casein and was removed when the latter was treated with ether. If centrifuged milk is used in the prepriation of the casein the amount of fat in the

other filtrate may be very small.



FIG 64 CALCIUM PROS-

8. Soponification of Butter. Dissolve a small amount of butter in alcohol made strongly alkaline with potasium hydroxide. Place the alcoholic potash solution in a casserole, add about 100 ml. of water and boil for 10 to 15 minutes or until the odor of alcohol cannot be detected. Place the casserole in a hood and neutralize the solution with suifuric

acid. Note the odor of volatile fatty acids, particularly butyric acid. Under certain conditions the odor of ethyl hutyrate may also be detected.

QUANTITATIVE ANALYSIS OF MILK"

1. Collection of Human Milk for Analysis. There are two methods of obtaining samples of breast milk for analysis

First Memor Express all the milk from one breast and mix thoroughlysecond Memor Draw one ounce of milk before nursing and one ounce after nursing Mix the two samples thoroughly. The best time for obtaining samples is 9 to 10 A.M.

2. Specific Gravity. This may be determined accurately by means of a pyknometer or more conveniently by means of a Soddet Vieth or Quevenue lactometer \(^1\) lactometer randing of 32° denotes a specific gravity of 1 032. The determination should be made at about 00°F (15°C) and the lactometer reading corrected by adding or subtracting 0 1° for every degree F above or below that temperature

3 Fat.

a BABCOCK'S CENTRIFUGAL METHOD PRINCIPLE This method involves the breaking of the emulson of fat in milk by means of concentrated sulfure acid centrifugation of the acid solution in the special tube shown in Fig. 65 and the subsequent reading of the percentage of fat in the graduated neck. Larger Babcock bottles are commonly

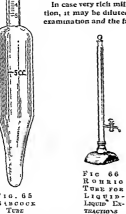
¹⁴ Official and Tentative Methods of Analysis 7th ed Washington D C Assoc of Official Agricultural Chemists 1950.

used for testing cow's milk or cream; these are supplied with pipets calibrated to deliver 18- or 9-g charges, respectively The method is accurate to within 0 5 per cent

Procedure. By means of a special narrow pipet, introduce milk into the tube up to the 5-ml. mark. Now add sufficient sulfuric acid (sp. gr. 1.83-1.834)

to fill the body of the tube and rotate the tube to secure a homogeneous acid-milk solution. Fill the neck of the tube with an acid-alcohol mixture.35 Centrifuge the tube and contents for one to two minutes and read off the percentage of fat by means of the graduated neck of the tube. If the top of the fat column is not at zero it may be brought there by the addition of hot water and a moment's recentrifugation.

In case very rich milk (over 5 per cent fat) is under examination, it may be diluted with an equal volume of water before examination and the fat percentage multiplied by 2. In the ex-



F1G. 65 Вавсоск



Fro 67 MOJONNIER. FLASK Designed to permit weighing, extraction, and decan tation without transfer of sample.

amination of cream it is customary to dilute the sample with four volumes of water and multiply the resultant fat value by 5,

b Roese-Gottlies Method Principle The milk is made alkaline and extracted repeatedly with petroleum benzin and the filtered extract evaporated to dryness in a tared flask. This method, together with the Babcock procedure, is "official" in the A O LC Book of Methods (see footoote 34, p 236) and is adaptable to butter, ice cream, dried milk, etc

Procedure. Transfer 10 g. of the sample to a Rohrig tube (Fig 66) or a similar apparatus (Mojonnier flasks* (Fig. 67) are widely used in dairy laboratories), add 1.25 ml of NH OH (2 ml. if the sample is sour), and mix thoroughly. Add 10 mi of 95 per cent alcohol and mix well. Add 25 ml of

²⁵ This mixture consists of equal volumes of amyl slookel and concentrated by drochloric

[&]quot; Obtained from Mojonnier Bros. Co , 4601 West Ohio St , Chicago 41 Ill

ether, shake vigorously for 30 seconds, add 25 mi of petrolcum benzin (redistilled slowly at a temperature below 65° C), and shake again for 30 seconds Let stand 20 minutes, or until the upper liquid is practically clear Draw off as much as possible of the ether-fat solution (usually 0 5 to 0 8 ml will be left) into a flask through a small, quick-acting filter Again extract the liquid remaining in the tube, this time with 15 ml of each solvent; shake vigorously 30 seconds after each addition and allow to acttle Draw off the clear solution through the small filter into the same llask as before and wash



FIO 68 SOXHLET LXTRACTION APPARATUS

the tip of the spigot, the funnel, and the filter with a few mi of a mixture of the two solvents, in equal parts, free from suspended II,0 To Insure complete removal of the fat, a third extraction is necessary (This third extraction yields less than I mg of fat if the previous solutions have been drawn off closely) Add a gliass bead and evaporate the ethers slowly on a warm surface, then dry the fat in a boiling water oven to constant weight Weigh the flask with a similar flask as a counterpolse Do not when the flask intendiately before weighing Remove the fat completely with petroleum benzin, beduct the weight of the dried flask with residue and bead to ubtain the weight of fat Finally, correct this weight by a blank determination on the reagents used

e SOXHET METHOD This classical procedure is suitable for the determination of fat in solid materials such as dried milk (Fig. 68)

4 Total Solids Introduce 2 to 5 g of milk into a weighed, flat-bottomed platinum dishⁿ (which may, if desired, contain 15 to 20 g of pure, dry sand spread over the bottom) and quickly ascertain this weight to millikrams. Expel the major portion of the water by heating the open dish on a water bath and continue the heating in an air bath or water oven at 97° to 100° C until the weight is constant (if platinum dishes are employed this residue may be used in the determination of ash according to the method described below.)

CALCULATION IS Divide the weight of the residue by the weight of milk used. The quotient multiplied by 100 gives the percentage of solids contained in the mdk examined.

5 Ath Heat the dry solids from 2 to 5 g of milk, obtained according to the method just given, over a very low fiame* until a white or light-gray ash is obtained II the determination is made directly on fluid milk, weigh quickly about 20 ml, add 6 ml concentrated HNO₁, evaporate, and ignite as above Cool the dish in a desiccator and weigh

6 Protein Introduce a known weight of milk (5 to 10 g) into a 500-mi Kjeldahi digestion flask and add 20 ml of concentrated sulfuric acid and

"The percentage of total solids may be calculated from the specific gravity and per centage of fat by means of the following formula

S = 025 L + 12 F

"Great care should be used in this ignition the dish at no time being heated above a faint redness since chlorides may volatilize

if Lead or aluminum fold dishes (which are much cheaper) make a very satisfactory substitute for the platinum dishes

S - total solids L - lactometer reading (third and fourth decimal places of specific gravity) P - fat content.

"Great rear about the content."

about 0.2 g. of copper suifate. Expel the major portion of the water by heating over a low flame, and finally use a full flame and allow the mixture to holl one to two hours. Complete the determination according to the directions eiven under the Kjeldahl Method, p. 874. If large amounts of milk are not available, a micro method may be used (see p. 880).

CALCULATION Multiply the total nitrogen content by the factor 6 3840 to obtain the protein content of the milk examined

7. Cosein. To 10.5 ml. (or a weighed equivalent) of fresh milk in a heaker, add 90 ml, of warm water (40° to 42° C.) and 1.5 ml, of 10 per cent acetic acid and stir. After 5 minutes' standing decant on an acid-washed filter, and wash the precipitate by decantation several times with cold water. Transfer the precipitate to the filter and repeat the washing twice. The filtrate should be clear or very nearly so. If the first portlons are not clear, repeat the filtration and complete the washing of the precipitate. Determine the nitrogen content in the washed precipitate and filter, as directed in Exp. 6.

CALCULATION Total N × 63S = cascia content of aliquot

8. Hort's Cosem Method Introduce 10.5 ml of milk into a 200-ml. Erienmeyer flask and add 75 ml of distliled water and 1.0 to 1.5 ml, of 10 per cent acetic acid.4 Mix the contents by giving the flash a vigorous rotary motion. The precipitated casein is now filtered off upon a 9- to 11-cm. filter paper. " Wash out the absorbed and loosely combined acetic acid by means of cold water. Continue the washing of both the casein on the filter and that adhering to the flask, until the wash water has reached a volume of at least 250 ml.

Now return the precipitate and paper to the original Erlenmeyer flash, add 75 to 80 ml, of neutral (carbon dioxide-free) water, 10 ml. of 0.1 N potassium hydroxide, and a few drops of phenolphthalein. Stopper the flask and shake it vigorously, hy hand or machine, until the casein has been brought into solution.42 Rinse the stopper with neutral (carbon dioxide-free) water and titrate the afkaline casein solution at once with 0.1 N hydrochloric acid until there is a disappearance of all red color,"

CALCULATION Subtract the corrected" acid reading from the 10 ml of alkali used The difference is the percentage of casein in the milk. For example, if it takes 67 ml of 01 N hydrochloric acid to titrate the alkaline solution to the end point and the check test was equivalent to 0 2 ml of 0 1 N acid, the casein value would be obtained as follows 10 - (67 + 02) = 31 per cent casein

Coogulable Profeso. Exactly neutralize the fitrate obtained under Exp. 7 with 10 per cent NaOH solution, add 0 3 ml, of 10 per cent acetic acld, and

¹⁰ The usual factor employed for the calculation of protein content of average or mixed foods from the nitrogen content is 6 25. It is based on the observation that proteins contain on the average 16 per cent nitrogen The factor 6 38 is used to calculate the protein content from the total mitrogen, since the total protein constituents of milk have a mean mitrogen content of 15 7 per cent

⁴⁴ In general 1.5 ml of acetic acid gives a clear solution which filters nicely, but occasion-

ally, when the milk has a low casem value at as advisable to use less acetic acid "The process of filtration may be retarded through the packing of the casein mass upon the filter paper. In this case conduct a fine stream of cold water against the upper point of contact of filter paper and casem By this means the casem precipitate is loosened and gathers in the apex of the filter This procedure is very essential. It is not necessary to re-

move the casem which adheres to the interior of the flask. 13 Solution is indicated by the disappearance of the white casein particles which would

otherwise settle to the bottom of the flask

^{44 \} check test should be run parallel with the entire determination. Even with special precautions as to neutrality it is generally found that an acid check of 0 2 to 0 3 ml will be obtained This check titration should be added to the volume of acid used in titration

heat on a steam bath until the protein is completely precipitated. Collect the precipitate on an acid washed filter, wash with cold 11,0, and determine the nitrogen as directed under Exp. 6

CALCULATION Multiply the total n trogen by the factor 6 38 to obtain the albumin and Llobul n content

10 Lactorse 4 To about 359 ml of water in a beaker add 20 g of milk, mli thoroughly, actdify the fluid with about 2 ml of 10 per cent acetle acid, and stir the acidified mixture continuously until a flocculent precipitate forms. At this point the reaction should be distinctly acid to litmus. Heat the solution to boling for one half hour, filter, runse the beaker thoroughly, and wash the precipitated proteins and the adherent fat with hot water Combine the filtrate and wash water and coocentrate the mixture to about 150 ml Cool the solution and diluted it o 200 ml in a volumetric flask. Thrate this sugar solution according to directions given under Benedict's Metbod (see p. 919).

Nyers recommends the following procedure for the determination of lactose in milk One part of milk is mixed with an equal volume of phos photungstic acid solution (70 0 g of acid and 200 ml of concentrated IJCI in I liter of water) and 2 to 3 parts of water Nix well, filter until clear, and titrate the clear filtrate against Benedict's solution (25 ml reduced by 67 mg of lactose)

Books Manuo for Lacrose in Milk Pipet 20 ml of mllk into a 100 ml volumetric flask From a pipet or buret add 12 ml of 10 per cent sodium tungstate and 12 ml of 25 is sulfured each Milk by rotating the flask, dilute to the mark-and filter Determine the lactose in the filtrate by titration with Benedict 8 solution

CALCULATION Make the calculation in the shove titration methods according to directions given under Benediet's Method (see p. 920) bearing in mind that 20 min of Benediet's solution are completely reduced by 0 067 g of lactose

If Micro Method for Lactore in Milk

I ancipus I actose is determined on the protein free filtrate of milk by a copper red ict nimit of following the method of Folm and Wu for blood sugar. This method is well suited to routine analyses.

Procedure Introduce 10 ml of milk into a 100 ml volumetric flask, add 2 ml of 10 per cent sodium tungstate Add gradually 2 ml of 25 N sulfure acid (or 16 ml of N/12 acid), mix well and let stand 5 minutes Diute to the mark with water and filter into a Folin Wu sugar tube introduce 1 ml of the filtrate and 1 ml of water into another tube place 2 ml of standard lactose solution 4 Add 2 ml of the Folin Wu alkaline copper solution (see p 568) to each tube, and heat in boiling water for 8 mlnutes. Gool and add diluted acid molybdate reagent (see p -688) to each tube After 1 mlnute, add diluted acid molybdate solution (1 4) to the 25 ml mark, mix, and compare in the colorimeter.

CALCULATION

المار

 $\frac{\text{Read ng of } \text{Standard}}{\text{Readn g of } \text{$\frac{\text{Standard}}{\text{Unknown}} \times 0.6 \times \frac{100}{0.01} \times \frac{1}{1000} = \text{Per cent lactose}}$

a. A method for lactose suitable for tissue analysis or where other sugars may be present in a gnificant a nounts in described by Malpress and Morrison Biochem J. 45 455, (1949). A star fact is bettern to dissolvent f. z. of lactoses in 0.2 per cent bear as as dark making up to a tissue of 100 ml. The working standard is prepared by diluting 3 ml of the stock soluti a to 100 ml with 0.2 per cent beanous and (2 ml = 0.6 ml).

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Epithelial and Connective Tissues: Bone and Teeth

EPITHELIAL TISSUE (KERATIN)

The major portion of hair, horn, hoof, feathers, nails, and the epidermal layer of the skin is made up of albuminoid proteins called keratins As a class the keratins are characterized by their extrume insolubility in the usual protein solvents, their lack of digestibility, and their high sulfur content most of which is in the form of cystine. These characteristics are not unrelated, the keratin molecule is considered to consist of closely packed polypeptide chains which are held together by the disulfide bond of cystine, the resistance to solvents and enzymes hein associated with the close packing of the chains. This view is supported by the fact that if wool, for example, is ground to a fine powder by mechanical means it hecomes more soluble and more digestible.

According to Block the vanous keratins may be further characterized as eukeratins and pseudokeratins. The eukeratins contain, in addition to other amino acids, the amino acids bistidine, lysine, and arginine in the approximate ratio of 1 4 12, with from 3 to 5 per cent sulfur, nearly all of which is cystine. They are found largely in the hair, nails, feathers, horn, etc. The pseudokeratins, found chiefly in the skin and nervous tissue contain less sulfur, from 1 to 3 per cent, and do not show the same instidine lysine arginine ratios as the cukeratins. It has been shown by ray crystallography that the intramolecular spatial arrangement of keratin can be modified under the influence of pressure or stretching. Such differences in molecular configuration of keratin may exist in the varied anatomic forms of epithelial tissue.

The keraturs are not acted upon by proteolytic enzymes. This is perhaps due more to lack of physical accessibility to the proteolytic enzymes than to lack of specific types of peptide linkages. Hence if the molecular structure of keratin is changed through chemical or mechanical means these enzymes digest the resultant product quite readily. The keratins appear to be digestible by an enzyme in the intestine of the clothes moth at pH 9 in the presence of the reducing medium yielded by the H₂S liberated through hacterial action in the intestine.

The composition of human hair is influenced by its color and by the race sex age and purity of breeding of the individual 1 It may be differ entiated from all other animal hair or wool by its high content of cystine

¹ Rutherford and Hawk J Bud Chem 3 4-9 (1907)

Most of the S of hair is in the form of cystine, with some cysteine usually present Human hair contains from 13 to 19 per cent of eystine, sheep's wool 8 to 14 per cent, and feathers 7 to 12 per cent It is claimed that the vitamius mositol, meotinie acid, pantotheme acid, and riboflavin are present in normal rat and human hair in about the same ratio as in other tissues 2

From the analyses of the skin of the dog, rabbit, and man for mineral constituents, it has been shown that there is an individual variability in composition in the same species as well as distinct differences among different species The addition of cystine to a diet deficient in this amino acid appears to favor the growth of hair to a greater extent than it affects the growth of hody tissue as a whole

Permanent waving is beheved to rest upon a change in the structure of the keratin molecule. The damp heat used in the waving process opens up the disulfide bonds of the keratin. If the hair he artificially curled while in this condition, then dried, the disulfide bonds again form and the wave is automatically held in position. On the other hand if the disul fide groups are changed to sulfhydryl groups by reduction the permanent wave may be made without resorting to heat. Hence the so-called "cold" wave

The addition of cystine to a sheep's diet will improve the weight and quality of the wool of the animal In rats, para aminohenzoic acid is also reported to have prevented the graying of hair Such findings on lower animals have not been successfully duplicated in controlled observations n human heings

Hyaluronic Acid and Hyaluronidase. So-called mucopolysaccharides seeur in epithelial and councetive tissues. The most important member of this group is hyaluronic acid. This may be hydrolyzed by the enzyme iyaluronidase & This enzyme was originally termed the spreading factor is it was thought to be importantly related to the spread of disease germs n the hody It seems that the hyaluronic acid present in the intercellular substance offers resistance to the spread of harmful microorganisms. However, many such microorganisms manufacture byaluronidase, which renders hyalurome acid more or less impotent, permitting the harmful germ to enter the tissues. Thus in rheumatic fever it has been claimed that the therapeutic use of salicylates inhibits the action of hyaluromidase The intact byaluronic acid then acts to prevent invasion of the tissues

In pure form, hyaluronidase which is also present in spermatozoa, leech heads, and the venom of snakes and bees, has been used clinically to facilitate the hypodermic administration of large volumes of fluids (hypodermoclysis) To assist the penetration of certain drugs into mucous membraues, to combat the formation of reual calculi to speed recovery from painful ankle sprains, and to aid in spreading the effect of local anesthetics over a larger area—these are other chinical functions ascribed to this enzyme. Evidence has accumulated which has east some doubt upon the supposed efficiency of hyaluronidase in hypodermoclysis 4

Novak and Bergeim J Biol Chem 155 283 (1944)
 Meyer I hysiol Ress 27 335 (1947)
 J Am Med 1480c 151 644 (1953)

EXPERIMENTS ON EPITHELIAL TISSUE (KERATIN)

Horn shavings or nail parings may be used in the experiments which follow

- 1 Millon & Reaction
- 2 Xonthoproteic Reaction
- 3 Glyoxylic Acid Reaction (Hopkins Cole)
- 4 Test for Cystine and Cysteine Sulfur

What amino acids do these tests show to be present in keratin?

CONNECTIVE TISSUE

T WHITE FIBROUS TISSLE

The principal solid constituent of white fibrous connective tissue is the albuminoid collagen This substance is also found in smaller percentage in cartilage bone and ligament but the collagen from the various sources is not identical in composition. In common with the keratins collagen is insoluble in the usual protein solvents presumably because of the close packing of the polypeptide chains as with the keratins It differs from keratin in containing much less sulfur Analyses show only 01 per cent cystine and 0 9 per cent methionine It contains no tryptophan and very little tyrosine It has been estimated that glycino represents one-third of all the amino acid residues in the collagen molecule, a second third of the molecule is composed of proline and hydroxyproline the other amino acids making up the remainder of the molecule. It is digested slowly by pepsin HCl but by trypsin only at temperatures above 40° C or after previous action of pepsin. One of the chief characteristics of collagen is the property of being converted by boiling acid or water to gelatin The process does not seem to be one of hydrolysis since there is no increase in amino nitrogen. Some intramolecular change may occur but the x ray diagram does not change This suggests that the alteration may be largely a physical one The amino acid composition is essentially that of gelatin (see p. 122)

In vitamin C (ascorbic acid) deficiency (scurvy) the intercellular cement substances are not deposited in a normal manner. It is helieved that in this condition there is a tendency to hemorrhage and to the slow heal ing of wounds because of the presence of abnormal collagen. In this con nection Robertson' reports no significant differences in the collagen con tent of skin liver lung kidney spleen costochondral junctions and teeth of normal and acutely and chronically scorbutic guinea pigs The same authors has shown that when guinea pigs in various nutritional states were fed glycine \13 the collagen isolated from liver lung muscle skin and bone contained \15 in varying concentrations The Lly cine isolated from the collagen of skin muscle and bone also contained an excess

Robertson J Biol Chem. 187 673 (19.0) * Ibid 197 490 (1902)

of the isotope. The concept of collagen disenses has received much attention. A wide variety of dissimilar diseases give definite evidence of fibrinoid changes ⁷ Hence the clinician has been warned that no special empliasis should be placed upon the occurrence of these changes and the clinical usefulness of the term "collagen disease" is questioned.

In the preparation of leather the collagen of the animal hide is purified and tanned. Certain precipitating agents such as tannic acid and the salts of heavy metals bring about the tanning.

The form of white fibrous tissue most satisfactory for general experiments is the tendo achillis of the ox. The fresh tissue has the following composition:

Water .	62 9 per cent
Solids	37 1
Inorganic matter	0 5
Organic matter	36 6
Fatty substance (ether-soluble)	1 0
Coagulable protein	0 2
Mucoid	1 3
Elastin	1 6
Collagen	31 6
Extractives, etc	0 9

The mucoid just mentioned is called tendomicoid and is a glycoprotein. It possesses properties similar to those of other connective-fussue mucoids, e.g., osseomucoid and chondromicoid.

Gelatin, the substance which results from the treatment of collagen with boiling water or boiling dilute acids, is sometimes classed as an albuminoid. It is probably better to consider gelatin a protein derivative not properly belonging to any of the recognized classes of proteins. Gelatin differs from the keratins and collagen in having a much simpler physical structure and in being easily soluble and digestible. In fact a large part of a gelatin solution can pass through membranes such as the walls of capillaries, because of its small molecular weight. Gelatin is nonantigenic and is not a complete protein from the nutritional point of view, since it is lacking in tryptophan and low or lacking in certain other amino acids (see p. 122). Thus it is not satisfactory as the sole dietary protein, but because of its ease of digestion and absorption it is used as an accessory protein in the diet, particularly in the case of convalescents. Attempts to remedy the dictary deficiencies of gelatin by supplementing the diet with the missing amino acids have not been successful; the reason is not known Gelatin gives a negative Hopkins-Cole test because it is lacking in tryptophan. The low content of tyrosine and cystine usually results in a negative or at the most a faintly positive reaction with Millon's reagent and the lead-blackening test. The isoelectric point of gelatin is about pH 4.7.

EXPERIMENTS ON WHITE FIBROUS TISSUE

The tende achilis of the ox may be taken as a satisfactory type of the white fibrous connective tissue.

⁷ J. Am Med. .1ssoc., 150, 220 (1952).

1 Preparation of Tendomucold Dissect away the fascia from about the tendon and cut the clean tendon into small pleces. Wash the pieces in running water, subjecting them to pressure in order to remove as much as possible of the soluble protein and lnorganic salts. This washing is very important Transfer the washed pleces of tendon to a flask and add 300 ml of haif saturated lime water. Shake the flash at Intervals for 24 hours Filter off the pieces of tendon and precipitate the mucold with dilute hydrochloric acid Allow the mucoid precipitate to settle, decant the supernatant fluid, and filter the remainder. Test the mucold as follows

a TEST FOR CYSTINE AND CYSTEINE SELFUR

- b Hydrolysis or Tendomucoid Place the remainder of the mucoid in a small beaker, add about 30 mi of water and 2 ml of dilute hydrochloric acid, and boil until the solution becomes dark brown Cool the solution, neutralize it with concentrated sodium carbonate, and test by Benedict's test
- 2 Collagen This substance is present in the tendon to the extent of about 32 per cent Therefore in making the following tests upon the pieces of tendon from which the mucoid, soluble protein, and lnorganic salts were removed in the last experiment, we may consider the tests as being made upon collagen
 - a BIURLT TEST
 - b. VANTHOPROTEIC REACTION
 - e GLYOXYLIC ACID REACTION (HOPEINS-COLE)
 - d Test for Cystine and Cysteine Sulfur Take a large piece of collagen in a test tubs and add about 5 ml of sodium hydroxide solution lieat until the collagen is partly decomposed, then add I to 2 drops of lead acetate and again heat to hound
 - e Formation or Gelatin from Collagen Transfer the remainder of the pieces of collagen to a casserole, fill the vessel about two thirds full of water, and boil for several hours, adding water at intervals as needed. By this means the collagen is transformed and gelatin is produced (see p. 245).
 - 3 Gelatin On the gelatin formed from the transformation of collagen in the above experiment (e), or on gelatin furnished by the instructor, make the following tests
 - a MILLON'S REACTION
 - h GLYOXYLIC ACID REACTION (HOPKINS-COLE)
 - C. TEST FOR CYSTINE AND CYSTEINS SULTING

Make the following tests upon a solution of gelatin in hot water

a Conducation Test Does it coagulate upon boiling?

b Precipitation by Alconol. Fill a test tube one half full of 95 per cent alcohol and pour in a small amount of concentrated gelatin solution. Do you get a precipitate? How would you prepare pure gelatin from the tendo achillis of the ox?

II YELLOW ELASTIC TISSUE (ELASTIN)

c ligamentum nuchae of the ox may be taken as a satisfactory type of the yellow elastic connective tissue. The principal solid constituent of this tissue is elastin a member of the albuminoid group. In common with the keratins and collagen elastin is an insoluble substance and gives the protein color reactions. It differs from keratin in amino acid composition and in the fact that it may be digested by proteolytic enzymes (see pp 122 and 183-184) Elastin is characterized by its low sulfur content and its high content of leuenie plus isoleucine

Yellow elastic tissue also contains mucoid and collagen but these are present in much smaller amount than in white fibrous tissue, as may be seen from the following percentage compositions of the fresh higamentum nuchae of the ox

Water	57 6 per cent
Solids	42 4
Inorganic matter	0 5
Organic matter	41 9
Fatty substance (ether colubic)	1 L
Coagulable protein	0 6
Mucoid	0 5
Elastin	31 7
Collagen	7 2
Extractives, etc	0.8

EXPERIMENTS ON ELASTIN

1 Preparation of Elastin (Richards and Gies) Cut the ligament into fine strips, run it through a meat chopper, and wash the finely divided material roold, running water for 24 to 48 hours. Add an excess of half-saturated fime water, and allow the hashed ligament to extract for 48 to 72 hours. Decant the lime water, where we all traces of alkali by washing in water, and then holf in water with repeated renewals until only traces of protein material can be detected in the wash water. Decant the fluid and boil the ligament in 10 per cent acetic acid for a few hours. Treat the pieces with 5 per cent hydrochloric acid at room temperature for a similar period, extract again in hot acetic acid and in cold hydrochloric acid. Wash out traces of acid by means of water and then thoroughly dehydrate by boiling alcohol and boiling ether in turn. Dry in an air bath and grind to a powder in a mortar.

- 1 Millon's Reaction
- 2 Youthoprotesc Reaction
- 3 Buret Test.
- 4 Glyoxylic Acid Reaction (Hopkins-Cole)
- 5 Test for Cystine and Cysteine Sulfur.

III CARTILAGE

The principal solid constituents of the matrix of cartilaginous tissue are chondromucoid, chondroalbumoid, and collagen Chondromucoid on decomposition yields protein and chondroitin sulfuric acid which has the following formula.

^a Later experiments (Half Reed and Tunbridge Asture 170 264 (1952)) show the presence of claudomucin a carbohydrate-containing protein The investigators was probably a mixture of clastin and elastomicin

Chondroitin sulfuric acid

On hydrolysis it loses sulfure and and forms chondroitin. The latter loses acetic and to form chondrosin Chrondrosin on hydrolysis yields chondrosimine CH-0H(CHOH)-CHN-HCHO and glucurome and COOH-(CHOH)-CHO Chondrosamine is apparently a galactosamine. The amino sugars are dealt with differently in metabolism from glucose or simple amino acids. They apparently do not form glycogen in the body Chondromucoids in various tissues differ from each other in the character of the protein only. Chondroitin sulfure acid differs from the sulfure acids found in the mucin of saliva, etc., in that the carbohydrate group in the latter is chitosamine (probably a glucosamine). It is of interest that chitin in the exoskelction of certain lower animals is a polysaccharide containing chitosamine and acetic acid. Chondroitin sulfure acid is found in tendomucoid and apparently also in osseomucoid, which are thus closely related to chondromucoid.

It has been shown that a deposition of radioactivity in the knee-joint cartilage of young rats follows the administration of radioactive sulfur in the form of sulfate. This radioactivity was concentrated in the chon droitin sulfate (chondroitin sulfate) and thyroxine influence the radioactivity pattern. For example in animals pretreated with thouracil the uptake of radioactive sulfur in the cartilage was reduced. The administration of thyroxine counteracted the action of thouracil. It was further shown through the use of isotopes that 0.134 g of chondroitin sulfate was formed per 100 g of ear tilage in 24 hours. The actual rate of sulfate formation was thus determined for the first time.

Chondroalbumoid is similar in some respects to elastin and keratin It differs from keratin in being digestible by proteolytic enzymes and in containing considerably less sulfur than any member of the keratin group It gives the usual protein color reactions

EXPERIMENTS ON CARTILAGE

1. Preporation of Chandroitus Suljutte Acid from Chandroitucoid. Free masal septa of cattle from bone and other extraneous material. Run 200 g. of the material through a meat chopper Add 400 ml. of 2 per cent NaOH Let stand for 2 days. Strain through cloth. Treat the residue again with 200 ml of NaOH for 2 days. Strain, Wash the residue once with water. Combine the extracts Acidify with acetic acid and then add an excess of barium carbonate Concentrate on the steam bath to half the volume Pour off the clear liquid Filter the remainder on a folded filter and add filtrate to the decanted solution Acidify Evaporate to about 80 ml. Centrifuge to remove protein and barium carbonate Drop the clear yellow liquid into 8 volumes of glacial acetic acid kept vigorously agitated (preferably with a turbine). Filter off the acid potassium salt with suction. Wash with glacial acetic acid and then with alcohol and ether.

Dissolve 8 g. of the product which still gives a slight hiuret test in 400 ml. of water, and while the solution is kept stirred drop in basic lead acetate solution to complete precipitation. Wash the precipitate several times by rubbing in a mortar with water and filtering with suction. Dissolve the precipitate in 10 per cent HCl. To the filtrate from the lead chloride add glacial acetic acid to precipitate all of the chondroitin sulfuric acid. Wash with glacial acetic acid, alcohol, and finally either

2. To Show the Presence of Sulfure Acid and of Reducing Sugars in Chandromucoid Treat 50 g of cartilage from nasal septum of the ox or cartilage rings from trachea of the ox with 100 ml of 1 per cent NaOH and let stand over night Pour off 50 ml of fluid, add 5 ml of concentrated HCl and boil for 30 minutes, bringling down to a low volume Toone portion add BaCl; solution and note the precipitate of BaSO. Neutralize another portion with sodium carbonate and apply Benedict's test to show the presence of a reducing carbohydrate group

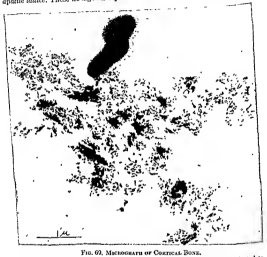
OSSEOUS TISSUE (BONE)

Bone contains from 14 to 14 per cent water, depending upon the type of bone and its location in the body. Of the fat-free dry matter of bone, from 30 to 50 per cent is organic, and the remainder morganic. Typical analyses of bone are given in Chapter 35 in connection with the discussion of rickets and vitamin D.

The organic portion of bone is similar in composition to cartilage. It contains collagen (ossein), osseoalbumoid, and osseomucoid. These proteins resemble the corresponding proteins in eartilage and tendon. Because of the collagen content, gelutin is formed when bone is holled with dilute and. The bone marrow also contains fat.

The morgame material of bone consists chiefly of calcium, phosphate, and carbonate with small amounts of magnesium, sodium, strontium, lead, citrate, fluoride, hydrovide, and sulfate These substances are present in amounts which correspond to 84 per cent Ca₂(PO₄)₂, 10 per cent CaCO₁, 2 per cent Ca₃(citrate)-, 10 per cent MgCO₁, and 20 per cent Na₂HPO₄ This composition is by no means constant Wide variations have been encountered For example, the molar proportions of PO₄ CO₂ can be anywhere from 1 8 to 3 4 These changes in composition are related to changes in the PO₄ CO₂ ratio of bloo

rum. The composition of blood serum in turn is influenced by nutrition, age, and disease. Despite the wide variations encountered in composition, x-ray diffraction studies reveal only one crystal structure, namely the apatite lattice. These hexagonal crystals in the bone are small in size, the



The diaphysis of a cat's femur autoclaved, agitated in blendor, and subjected to ultrasonies. The two bacteria are Bacillus aerogenes and give an idea of the relative size of the crystals in relation to bacteria. Magnification 34,500.

Courter, R. A. Bolsson and F. W. Babo, and Surner.

dimensions seen in the electron microscope being about 500 \times 250 \times 100 Angstrom units (Fig. 69).

Because of this small size, the bone crystal offers a large surface that can adsorb other compounds than those which account for the crystal lattice itself. Evidence for this theory was the preferential solubility of carbonates and citrates in bone. There are many alternative theories, but the one which appears to be most acceptable is that of Hendricks, who visualizes the crystal structure of bone salts as due to hydroxyapatite

Sobel, Rockenmacher, and Kramer J Bust Chem., 158, 475, and 159, 159 (1945).

Chap. 9

Ca10(PO4)6(OH)2, in which F may replace OH, with the rest of the composition held by surface forces (Fig. 70).

Despite the possible variation in bone composition indicated in the discussion above, under some conditions the inorganic material of bone is rather constant in composition. Observations on the constancy of composition of bone during fasting are of interest in this connection. The

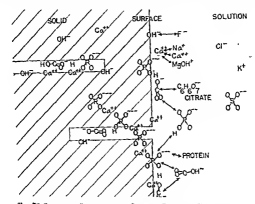


Fig 70 Schematic Indications of Surface Compound Formation in THE BONE SALT.

Courtesy, S B Hendricks, "Comments on Crystal Chemistry of Bone" from the Fourth Conference on Metabolic Interrolations of the Joseph Macy, Jr Foundation.

percentage composition of the dry fat-free femure of two dogs, after the animals had fasted for 104 and 14 days respectively, was as follows:

Dog No.	Length of Fast	Ash	N	Ca	Mg	P
1	104 days	61 5	46	23 3	0 5	12 8
2	14 days	61 7	41	23 2	0 5	12 9

The marked uniformity in composition notwithstanding the wide variation in the fasting periods is significant. The tensile strength of the femur of the dog has been found to be at least 25,000 pounds to the square inch whereas that of oak is 10,000 and that of cast iron 20,000 pounds to the square inch.

The percentage compositions of normal human bone and of hone from a case of osteomalacia with respect to certain elements are given in the

following table:

0	Kind of Bone			
Constituent	Normal	Osteomalacia		
Calcium	20 2	10 8		
Magnesum	0.1	03		
Hosphorus	8.6	o 3		
Silfur	0 1	0.6		

Hammett has shown in the case of albino rats that there is an increase in calcium and a decrease in magnesium and phosphorus during the first 75 days of growth. Moreover the bones of the female have a higher calcium content than the bones of the male.

Chemistry of Ossification. Evidence from many sources indicates that there is an active metabolism in bone. The mineral matter of bone may be drawn upon in case of need elsewhere, as in the formation of milk hy the lactating animal and in the formation of the hones of the fetus during pregnancy This utilization of hone material should be regarded as a resorption of hone rather than a simple decalcification since both the organic and inorganic components of bone disappear during the process Studies with radioactive phosphorus as a tracer 10 have shown that almost immediately after the introduction of radioactive phosphorus into the blood there is an appreciable uptake of the labeled phosphorus by the bones, indicating a turnover or metabolic interchange hetween the bone and plasma or lymph phosphorus During a 50-day period in the life of the rahbit, for example, half the scapula may he replaced, with a 30 per cent turnover of the epiphyseal phosphorus, diaphyseal hone metaholism appears to be considerably slower There are indications of a labile fraction of bone which is in equilibrium with the plasma and a stable fraction which is in equilibrium with the labile portion

The mechanism of calcification is not clear. That the solubility product principle operates is indicated from studies of healing rickets in vivo as well as in vitro. Not until the product of the ionic concentration of cal cium and phosphate exceeds a minimum is there any new mineralization in the embryonic or rachitic bone cartilage matrix. This minimum is not

the same for all bones, as is shown in the table below 11

THE SOLUBILITY PRODUCT PRINCIPLE AND NEW CALCIFICATION

Bone	Ca	×	P.
Fmbryome		16	_

Thus one must postulate the existence of local factors in the calcifying site whose concentration determines the level at which the solubility product operates Several explanations have been offered for the nature of this local factor Phosphatasc, present in bone cartilage, has been suggested as the enzyme which on splitting organic phosphate causes supersaturation of calcium phosphate, thus promoting precipitation It has also been suggested that glycogen and the enzymes responsible for phosphorylative glycogenolysis provide both the organic phosphate and the local mechanism More recently, however, Sobel advanced the view that a mucopolysaccharide, chondroitin sulfate, is the local factor, based on (1) the presence of this compound at the site of mineralization, (2) the observation that when phosphatuse and enzymes of the phosphorylative gly cogenolysis are destroyed by heating and other measures, bone sections will still mineralize, (3) studies of the reversible mactivation of the ability of bone to calcify in vitro, by cations such as Be++ and Cu++ as mactivators and Ca++ as the reactivating ion

Mthough the precipitate found in bone is composed of [Ca₂(PO₄)₂]_a CaCO₃ (n = 1 8 to 3 4), studies of the physicochemical mechanism indicate that the first aggregate formed is CaHPO₄ which is rapidly transformed to hydrovapatite So far however, no direct evidence has been found for the existence of CaHPO₄ in bone

Vitamin D promotes calcification especially when added to rachitogenic diets Under the influence of the antirachitic vitamin, the scrum Ca × P product becomes elevated as shown below

INFLUENCE OF VITAMIN D ON SERUM CA AND P*
(Mean Values)

Diet per cent		Serum mg per cent			
Ca	P	Са	P	Cat	P†
1 20 0 19 0 03	0 12 0 12 0 76	11 7 9 4 5 6	2 1 4 7 7 5	13 3 11 1 8 8	3 4 6 0 8 4

 ²³⁻day-old rats on a rachitogenic diet for 30 days
 Receiving 100 units of vitainin D daily

This clevation in serum $Ca \times P$ is probably the main reason for the effectiveness of vitamin D. In diseases like Fancon's nekets in which this elevation of serum $Ca \times P$ does not take place, the rachitic condition persists. Though the mode of action of vitamin D is not fully clear, there is both increase in the absorption of calcium from the intestinal tract and increased resorption of phosphate from the kidney. Both of these effects of vitamin D would tend to elevate serum $Ca \times P$ and thus promote calcification. The finding of a high phosphatase content of the blood in ostetits deformans, nekets, and other generalized bone disorders, and a decrease of phosphatase in the tissues of the animals red a det high in vitamin D, are cfiniterest in this connection Calcium metabolism and

P = inorganic phosphate

deposition are also influenced by the endocrine glands (see Chapter 26, Hormones)

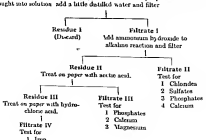
EXPERIMENTS ON OSSEOUS TISSUE

- I Decalcification of Bone Treat a strip of rib hone with dilute IINO₁ and permit it to soak for 6 to 8 days, renewing the acid every 1 to 2 days. After complete decalcification, wash the bone, split it, and remove the marrow. Now cut the bone into small pieces, wash it to remove acid, and boil the material in water for at least one hour Filter, concentrate the filtrate, and permit it to cool. What is the characteristic of the cooled filtrate? What has been produced from the bone?
- 2 Qualitative Analysis of Bone Ash Take I g. of bone ash in a small beaker and add a little dilute nitric acid. What does the effervescence indicate? Stir thoroughly, and when the major portion of the ash is dissolved add an equal volume of water and filter. To the acid filtrate add ammonlum bydroxide to alkaline reaction. A heavy white precipitate of phosphates results. (What phosphates are precipitated here by the ammonia?) Filter and test the filtrate for chlorides, sulfates, phosphates, and calcium Add dliute acetic acid to the precipitate on the paper and test a little of this filtrate for calcium and phosphates Heat the remainder of the filtrate to bolling and add (NII.):CO; and NII,Cl slowly to this hot solution as long as a precipitate forms. Fifter off the precipitate of CaCO, and wash with hot water until free from alkali.12 To the filtrate add a solution of NasiiPO, make strongly alkaline with NILOII, and note the formation of a white precipitate of ammonium magnesium phos phate (MI, MgPO.) Examine the crystals under the microscope and compare with those shown in Fig 226 To the precipitate on the filter paper, which was Insoluble in acetic acid, add a little dilute hydrochloric acid and test this last filtrate for phosphates and Iron

Reference to the scheme below may facilitate the analysis.

Bone Ash

Add dilute nitne acid stir thoroughly, and after the major portion of the ash has been brought into solution add a little distilled water and fifter



EPITHELIAL AND CONNECTIVE TISSUES BONE AND TEETH

MICROESTIMATION OF THE INORGANIC CONSTITUENTS OF BONE 13 Sobel et al, have described a scheme for analysis of CO2, Ca, P, and total base which is serial in operation and requires only one weighed specimen. The complete analysis can be performed on as little as 5 mg of sample.

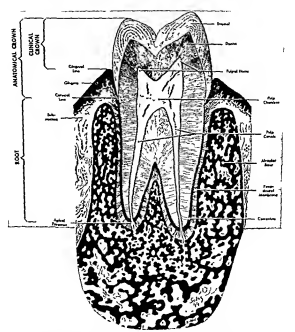


Fig 71 Schematic Drawing of a Tooth and the Surrounding Structure Courtesy, Zeisz and Nuckolls Denial Anatomy St. Louis C V Visby Co 1949 p 20.

TEETH

Composition. Teeth are composed of four tissues enamel, dentin, and cementum which are highly calcified, and the dental pulp which is uncalcified and is surrounded by the dentin (Fig. 71). Enamel covers the

¹² Sobel Rockenmacher, and Kramer J. Bud. Chem. 152, 255 (1944)

dentin in the crown of the tooth. It is the hardest substance in the body and contains the smallest amount of water. Cementum contains about 30 per cent organic matter. It covers the dentin in the root of the tooth, which is contained in the supporting structures known as the periodontal tissues. These tissues, which include the gingiva and alveolar hone, hold the teeth in line. The periodontal membrane, which is adjacent to the cementum, connects the teeth to the gingiva and alveolar bone.

The average composition of human enamel and dentin is shown in the following table

COMPOSITION OF HEMAN ENAMEL AND DENTIN

	Enamel	Deniin	
Constituent	Per Cent on Dry Busis*		
Asb	97	72	
Calcium	35 8	26 5	
Magnesium	04	0.8	
Sodium	07	02	
Potassium	03	0 07	
Phosphorus	17 4	12 7	
Carbon dioxide (from carbonate)	28	3 06	
Chlorine	03	0 03	
Fluorine	0 0112	0 0204	
Iron	0 0218	0 0072	
Organic matter	1	20	

Water which is not femoved at about 100°C the temperature to which the material is usually leased before analysis was included in the weight from which calculations with made. The percentage of such combined water is approximately 2 for enamed and 8 for denian.

The average values for the composition of whole buman teeth are shown in the following table

Il hole Human Teeth

On the hasis of chemical analysis, it was thought at one time that most of the calcium and phosphorus in the dental tissues existed as $\mathrm{Ca}_3(\mathrm{PO}_4)_2$, a view formerly held also for hone. However, x-ray crystal-lographic studies suggest that the crystal structure of most of the calcium phosphate in the dental tissues is similar to that of hydroxyapatite, $\mathrm{Ca}_{10}(\mathrm{PO}_4)_{\mathrm{e}}^*(\mathrm{OH})_2$. The striking similarity of x-ray diffraction photographs of hydroxyapatite and cnamel is shown in Fig. 72, which contains also the photographs of fluoapatite and chloroapatite. There is some evidence which suggests that in the case of dentin and ementum additional phosphorus in the form of either PO_4^- or HPO_4^- or some other ion involving phosphorus is occluded in the hydroxyapatite molecule. The apatite salt

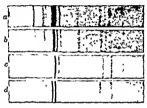


Fig. 72. X-rat-Differaction Photographs
(a) Human dental enamel, (b) hydroxyapatite, (c) fluoapatite, (d) chloroapatite
From Theula, Glock and Murray: Trons. Faroday
Soc. 35, 358 (1939)

nown as tricalcium phosphate hydrate, which has been claimed by some o be the main morganic salt in dentin and cementum, is probably hydroxyapatite with occluded phosphate ion. It is claimed that smaller imounts of carbonateapatite, chloroapatite, and fluoapatite are present nenamel, and that dentin contains some fluoapatite. A portion of the oddium, potassium, and magnesium in the dential tissues may be part of the patite lattice by substitution for part of the calcium. Enamel and lentin on spectrographic examination have heen found to contain numerous trace elements, including aluminum, harium, boron, chromium, copper, lead, lithium, manganese, nickel, silicon, silver, strontium, tin, titanium, vanadium, and zine.

The electron microscope as well as x-ray studies reveal that the crys-

tals of enamel are larger than those of hone or dentine.

The composition of growing teeth can be influenced in a predictable manner, high-carbonate teeth being produced on a high-calcium, low-phosphate dict and low-carbonate teeth on a low-calcium, high-phosphate dict. Evidence to date indicates that high-carbonate teeth are more susceptible to caries. 16

Posner and Stephenson: J. Denial Research, 31, 371 (1952).

Sobel and Hanok; J. Biol. Chem., 176, 1103 (1948).
 Sobel. Fourth Janah Macy Conference on Metabolic Interrelations, 1952, p. 261.

The organic matter of enamel consists mainly of the protein keratin which on the basis of the proportion of histidine lysine, and arguine is of eukeratin nature (see p. 184). Smill amounts of cholesterol and phopholipide are all o present. The albuminoids collagen and elastin are found in dentin, collagen representing the major organic constituent of this tissue. Glycoprotein is present in dentin along with small amounts of cholesterol and phopholipide. Collagen is the main organic constituent of cementum. Citrate has been identified as a constituent of human teeth, we to eight times more being present in dentin than in enamel.

Chronolour or Heman Dentition (Logan and Aronfeld slightly modified by McCall and Schour)

Tooth		Fırst eviderce of calcification	F namel complete I	Root completed
dentiti n Lateral incisor Cusp d First m lar		Lateral meisor 4½ mos mutero Cusp d 5 mos in utero First m lar 5 mos in utero		114 2 yrs 112 2 yrs. 212 3 yrs 2 214 yrs 3 yrs
Permanent dentition Upper jaw	Central incesor Lateral meisor Cuspid First I icuspid Second bicuspic First molar Second molar Third molar	3 1 mos 1 yr 4 J nios 1½ 1¾ yrs 2 2½ yrs. At 1 nrth 2½ 3 yrs 7 9 yrs	4-5) rs 4 3) rs 6-7 yrs 5-6) rs 6 7) rs 2 1 ₂ 3) rs 7 8) rs 12-16 yrs	10 yrs 11 yrs 13 lo yrs 12 13 yrs 12 14 yrs 9-10 yrs 14-16 yrs 18-20 yrs
Lower 12w	Central meisor Lateral meisor Cusj id First b euspi I Second bieusp First molar Second molar Third molar	3-4 mos 4-5 mos 13/4 2 yrs	4-0 yrs 4-0 yrs 6-7 yrs 5-6 yrs 6-7 yrs 21 2 3 yrs 7 8 yrs 12-16 yrs	9 yrs 10 yrs 12 14 yrs 12-13 yrs 13-14 yrs 9-10 yrs 14-15 yrs 18-20 yrs

Several investigations have been made in an attempt to ascertain whether differences in chimal composition exist between enamel of sound teeth and the intact enamel of carous teeth \(^{\infty}\) osgnificant differences were found for the contents of calcium, magnesium, phosphorus, and carbonate \(^{\infty}\) multi studies on dentin have likewise given negative results On the other hand it has been reported? that enamel from sound teeth centains significantly more fluorine than does the sound enamel from earnous teeth although no difference was found for danti However.

¹⁷ Armstrong and Brekh is J Dental Research, 17 333 (1938)

this difference between sound and carrous teeth was not found in all localities where these determinations were made 18 (See p. 262 for additional data on the relationship of fluorine and teeth) No difference was found in the ash, calcium, and phosphorus contents of the root dentin of teeth from pregnant aud nonpregnant women, this fact suggests that there is no basis for the view that minerals are withdrawn from the teeth during pregnancy Furthermore, chinical investigation has shown that the incidence of dental decay in pregnant women is no greater than in nonpregnant women of corresponding age

Chronology of Dentition. The ages at which the deciduous and permanent teeth begin to ealcify and are completed are shown in the table19 on p 258

In a study of calcification of teeth at birth, it was found that all the teeth of both jaws contained approximately 212 mg of calcium and 100 mg of phosphorus At the time of birth the lateral incisors, the cuspids, the first deciduous molars, the second deciduous molars, and the first permanent molars contained respectively 20, 7 5, 21, 11, and 0 1 per cent of the calcium present in the fully erupted teeth. These data thus indicate that the postnatal period is of far greater importance than the prenatal period in the calcification of the deciduous as well as of the permanent teeth

SYSTEMIC EFFECTS

In studies dealing with the response of the dental tissues to systemic influences it is important to distinguish between the developing and the adult teeth, and between the deutal tissues of bmited growth, as in humans, and those of continuous growth like the incisors of rats A great deal of confusion has been caused by failure to recognize this distinction

Adult teeth of hmited growth are probably only slightly affected by systemic influences. That such effects may occur is suggested by studies in which radioactive isotopes have been used. For example, it was shown that following the ingestion or injection of sodium phosphate containing radioactive phosphorus, small amounts of the isotope appeared in the enamel and dentin, but much smaller amounts were present in the former than in the latter However, the slight extent to which this occurs is shown by the fact that after the ingestion of 900 mg of sodium phosphate containing isotopie P by an individual twenty-five years of age, about 1/300,000 of the labeled P entered a single tooth On the basis of such data it was calculated that the replacement of 1 per cent of the total tooth phosphorus by that taken up in food would take about 250 days

For the proper calcification of the teeth the diet must contain adequate quantities of calcium and phosphorus and eertain vitamins. Some of the hormones likewise play a fundamental role

Vitamins, Calcium, and Phosphorus. The presence of vitamin A is essential to the process of tooth formation. Dietary deficiency of this vitamin during the period that the teeth are undergoing development

⁴ McClure J Dental Research 27, 287 (1943)
4 Logan and Kronfeld Hutopothology of the Teeth and Their Surro inding Structures revised and re-edited by Boy to Fhiladelphia Lea & Februer, 1949 p. 41

causes the ameloblasts (enamel forming cells) to atrophy, resulting in hypoplastic (incompletely developed) enamel which is imperfectly calcified. There is also atrophy of the odontoblasts (dentin-forming cells) so that the dentin laid down is also incompletely calcified. It is claimed that the primary effect is on the enamel, irregularities in the development of the dentin being secondary to the chamel hypoplasia. Deficiency of vitamin A in rats has been found to produce an increase in the percentage of magnesium in the incisors in spite of the fact that the ash content is decreased. A characteristic effect of vitamin A deficiency—namely, the substitution of stratified keratinizing epithelium for normal epithelium—has been observed in the gingiva. Malformation of alveolar hone has been produced in dogs by feeding them vitamin A-deficient diets.

Vitamin C is also important for the functional activity of the formative cells. Dietary deficiency in vitamin C during the period of tooth development causes impairment of the odontohlasts, leading to defective calcification of the dentin accompanied by hemorrhagic and degenerative lesions in the pulp tissue. The changes in the dentin and pulp have been attributed to an inability to produce and maintain intercellular substance. Defective enamel formation, which has also been observed, is said to be secondary to retarded dentin deposition. Pathological changes in gingival tissue have been demonstrated to be due to deficiency of vitamin. C in the diet. These conditions, as well as those in the teeth themselves, have been shown to respond favorably to the addition to the diet of

vitamin C

The degree of mineralization of the teeth depends on the relative as well as on the absolute quantities of calcium and phosphorus in the diet, and also on the amount of vitamin D or exposure to ultraviolet rays. Even with optimal quantities of calcium and phosphorus the presence of vitamin D is essential to the development of perfect teeth Hypoplastic defects in the enamel and imperfect calcufaction of the dentin have been found in neckets in children. It has been reported that an increase in the vitamin D intake brings about a reduction in the incidence of dential decay in children. Similar results have been produced in rats by adding vitamin D to a caries-producing diet.

Although both teeth and bones are highly calcified tissues, metabolic effects in the one do not necessarily parallel those in the other. For example, although the percentage of ash of the incisor teeth of rats fed a low-calcium, low-phosphorus det was lower than that of animals on a normal dict, the absolute amount of ash on the deficient diet increased about 15 per cent during a period of nine weeks. In the normal animals there was as increase of 77 per cent. But during this period there was a reduction of 16 per cent in the absolute amount of ash in the femirs of the animals on the deficient diet, while in the normal animals there was an increase of 33 per cent. Thus there was a continuous deposition of minicrals in the incisors of the rats on the deficient diet, while bone underwind demineralization. Furthermore although the percentage of ash, calcium, and phosphorus of the tibiae of rats may be reduced to about the same extent on a high-calcium, low-phosphorus richitic diet, and on a low-calcium high-pho-phorus richitic diet the values for these constitutions.

uents are unaffected in the meisors of the rats on the high calcium diet, but are somewhat reduced on the low calcium diet. These facts indicate that in some fundamental way there is a marked difference between the mineral metabolism of teeth and bones.

Hormones Normal tooth growth requires an adequate supply of thyronine Hypothyroidism in children causes retriction in dental development although no structural defects of the teeth have been detected in this condition (Structural defects and impaired calcification have, however, heen found after thy roidectomy in monkeys and rabbits) Reduced activity in the thyroid gland during morphodifferentiation, that is, in utero or in the first year of ble, results in a reduction of the size and an alteration in the form of the crowns of the teeth. The size and form of the root may be altered by disturbances in the thyroid gland in later periods. In hypothyroidism there is delayed resorption and shedding of the deciduous teeth, whereas in hyperthyroidism which is relatively rare in children there is early shedding of the deciduous teeth and accelerated inition.

Deficient parathyroid function may interfere with calcium and phosonous metabolism to such an extent that growing teeth fail to calcify operly. If this occurs in infancy, enamed hypoplasia and disturbances the calcification of dentin occur in the tissues calcifying at that time involved in the animal hypoplasm in rats, the incisor teeth, which grow moughout life in this animal, become brittle and distorted, whereas fully rimed enamed and dentin show no changes. Administration of paraisormone to parathyroidectomized rats results in normal calcification of ce enamed and dentin calcifying during the period of treatment, but does it restore to normal the structures that were formed in the absence of arithyroid function. Inhyperparathyroidsm although calcium and phosonous are withdrawn from hone there is no such withdrawal from the oth. The teeth may become loose as a result of resorption of alveolar one, but the teeth remain well calcified in spite of the severe drain upon one.

The pituitary gland plays an important part in the rate of development of the teeth. Dysfunction of this gland usually does not occur before to 6 years of age, that is, before the crowns of the permanent teeth (expt the second and third molars) are fully formed. Since cruption of the zeth is retarded, the chineal crown may be smaller than normal. However, the size of the anatomic crown is not affected. Juvenile dental characteristics persist as shown by large pulps and incompletely developed dots. Disturbances occur in the calculation of the dentin and alveolar one. One of the main dental effects in hyperpituitarism is an acceleration a the rate of cruption of the teeth.

In experimental hypogonadism in the male rat, evidence of disturbed alcification was apparent 60 to 90 days after castration. The male horsone therefore appears to have an effect ou the calcification of teeth in hypergonadism in humans, growth, calcification, and eruption of the each are accelerated, but not to the same extent as is skeletal growth.

Disturbances of the calcification of growing dentin have been reported

n experimental adrenalectomy in the rat

FIUORINE AND IEEFII

The mass occurrence of a characteristic dental defect known as mottled enamel or dental fluoro is was observed about 1908 in the inhabitants of Colorado Springs Approximately 80 per cent of those who were born and raised in this community were affected to some degree. Those who came to the locality after their teeth had been completely formed were not afflicted with mottled enamel after residence there Surveys of other communities revealed a similar dental defect. Analysis of the water in the communities where mottled enamel was found revealed the presence of excessive amounts of fluoride. At Bauxite, Arkansas for example, the fluoride content of the water was 13 to 14 parts per million (13 to 11 mg per liter) a condition which was subsequently remedied

Enamel fluorosis is limited to children in whom the cnamel of the per manent teeth has not been completed Enamel formation is completed except for the third molar, during the seventh to eighth year. It is completed in the third molars between the twelfth and sixteenth years. The disease is characterized by chalky white patches distributed over the en amel, which is pitted and corroded in extreme cases resulting in loss of the normal translucency The fluorine content of mottled enamel may be two to three times that of normal enamel the fluorine content of dentine is likewise increased when water containing excessive amounts of fluoride

is ingested during infancy and childhood

It was found that in communities where dental fluorosis existed there was less dental decay than in comparable communities where the water was practically fluoride-free In one town where the water supply con tained 18 parts of fluoride per million parts of water there was about one-third as much dental decay as in a comparable group of children in another town where the water was almost fluoride-free About of per cent of the children who were raised in the former community showed a mild dental fluorosis On the basis of this early evidence and sub-equent studies many cities and towns are now adding fluoride to the water supply in an effort to reduce the meidence of dental caries Fluoride is added to bring the concentration up to approximately 1 part per million, a level at which dental fluorosis is avoided From the favorable reports regarding reduction of dental caries following this procedure the addition of fluoride to water to the level of 1 part per million appears to be a worthwhile public health measure

Another effort to reduce dental decay has my olved topical application (application directly to the teeth) of from 0 1 to 2 per cent solutions of sodium or potassium fluoride According to several reports20 the incidence of dental caries has been materially reduced by this procedure other reports have been negative. However the evidence in favor of topical application of fluoride solutions appears to outweigh that against it. In this connection it is of interest to note that topical application of a 2 per cent sodium fluoride solution to the teeth of dogs has been found to increase

Knudson and Armstrong Am. J Public Health 14, 239 (1944) Klinkenberg and B bby Dental Research 29 4 (19.0)

the fluoride content of the teeth to a significant degree.²¹ Furthermore, enamel which has been treated with fluoride solutions is less soluble in acid than untreated enamel,²² a fact of importance because in the process of dental decay, the mineral salts of enamel, and later of the dentin, are dissolved by acids which form as a result of the action of enzymes on carbohydrate.

EXPERIMENTS ON TEETH

Anolysis of Teeth. Place a tooth in 25 ml. of dilute nitric acid and allow to stand over night. On the solution abtained run an analysis for inorganic elements according to the procedure given for bone ash (p. 254). Separate analyses may be made on dentin and enamei. Note the very slight amount of organic matter in enamel.

Quontitative Analysis of Teeth. Weigh a clean dry tooth on an analytical balance. Put in a beaker, add 25 ml. of dilute HCl, and let stand over night. Dilute to exactly 100 ml. and filter.

Determination of Colcium. Pipet 10-ml. portions of tooth solution into each of two 250-ml. beakers. Add to each 20 mi. of 2,5 per cent oxain acid solution, 2 drops of methyl red (saturated alcoholic solution), about 70 mi. of distilled water, and then concentrated NH,OH drop by drop with vigorous stirring. When a precipitate of calcium oxaiate begins to form add more ammonia very slowly until the color changes to an intermediate color between red and yellow. Let stand over night. Complete the determination according to directions in Chapter 31. Calculate the percentage of Ca in the tooth.

Determination of Phasphorus. Pipet 10 ml. of tooth solution into a 150-ml. beaker, add 2 drops of phenolphthalein solution and NaOH until the color just turns faint pink. Carry out a uranium acetate titration or determine colorimetrically according to the directions given in Chapter 31. Calculate the percentage of P.

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¹¹ Hord and Lilis J Dental Research, 39, 360 (1951)

[&]quot;Sucse and I osdick J Dental Research, 30, 177 (1951).

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10

Muscular Tissue

COMPOSITION OF MUSCLE

Muscular tissue constitutes about 40 per cent of the body weight, and is, therefore, the largest single tissue component of the hody There is great variability in the physiological and finer morphological characteristics of individual muscles but it is usual to classify them as striated (skeletal, voluntary), cardiac, and smooth muscles Chemically, striated muscle has been investigated more extensively than cardiac and, especially, smooth muscle

In round numbers, striated muscle consists of 75 per cent water, 20 per cent proteins, and 5 per cent other solids, such as morganic salts and the

so-called "extractives"

EXTRACTIVES OF MUSCLE

Under the name extractnes are classed a number of muscle constituents which occur in small amounts in the tissue and may he extracted by water, alcohol, or ether There are two classes of these extractives, the non natrogenous and the nitrogenous Grouped under the non-nitrogenous compounds are glycogen, hexosephosphates, lactic acid, mositol, and fat In the class of nitrogenous extractives are creatine phosphate, purme bases (\anthine, hypo\anthine), uric acid, adenylic acid, mosmic acid, carnosine (ignotine), anseriue, and carmtine (novaine) All of these substances, and many more, have been obtained from dead muscle, there is evidence that some of them are produced largely by post-mortem reactions and are not present as such in significant amounts in living muscle Other extractives hesides those enumerated above have heen described, and there are undoubtedly still others whose presence remains undetermined A detailed consideration would, however, he unprofitable in this place

NON-NITROGENOUS EXTRACTIVES

Lipides. Like most organs, muscles contain storage fat (not dissimilar to that in other depots) as well as essential lipides Phospholipides predominate among the latter It has been estimated that mammalian muscles contain on the average 4.5 per ceut phospholipides and 0.25 per ceut cholesterol, in terms of dry weight For the heart, these proportions are 7 5 and 0 5 per cent, and for smooth musele 3 and 0 75 per ceut, respectively (Bloor)

Glycogen. The chief carhohydrate of muscle is the polysaccharide glycogen (see p 87). It may he prepared from muscle or other tissue hy

extraction with boiling water followed by precipitation with alcohol By such methods, opalescent solutions of amorphous glytogen are obtained but by more careful procedures it can be shown that in the living tissue, glytogen occurs in the form of submicroscopic granules, comparable per haps to vegetable starch grains, although much smaller

The glycogen content of muscle varies and is reduced by intense activity, but is not, as is liver glycogen, significantly depleted as a result of a demand for earbohydrate elsewhere in the body Glycogen occurs also in the heart, its behavior in this organ is not the same as in skeletal mus-

ele, and requires further study

Hexosephosphates, Of the several existing hexosephosphates, resting muscle contains only the so-called Emhden ester. This is an equilibrium mixture of glucose-6-phosphate and fructose-6-phosphate (see p. 273), the only intermediates of the glycolytic cycle which accumulate to an appropriable extent under normal conditions.

Inositol occurs in muscle as it does in many tissues, but its mode of

combination and its function are unknown (see Chapter 35)

Lactic Acid. This substance occurs in muscle in variable quantity, dependent on the physiological state of the organ Resting muscle, after careful extraction, contains only very small amounts of lactic acid (e.g., 0.015 per cent in frog muscle) but it is found in larger amount after stimulation, in anaerobiosis, and post mortem. Rigor mortis is associated with lactic acid production and with depliction of adenosinetriphosphate

MITROGENOUS EXTRACTIVES

Creatine. Creatine, methylguanidinoacetic acid, is found in varying

amounts in muscles of vertebrates, frequently representing about 0.3 to 0.5 per cent of the wet tissue. The creatine of the body is maintained by synthesis of glycocyamine and its methylation and a regular excretion of creatine takes place in the form of its anhydride creatinine, in the unine, dependent somehow on the development of the muscular tissue. In muscular dystrophies, whether spontaneous as in man or induced in animals by vitamin E deficiency, this creatine excretion is greatly enhanced.

Creatine forms colorless monoclimic prisms which are tasteless to some individuals but butter to others and which decompose with marked efforcescence at about 291? C (corr.) It is soluble in warm water and practically insoluble in alcohol and ether Upon boiling a solution of creatining in the processing of the processing of the processing in the processing of the proces

Phosphocreatine. Though creatine is present in meat extracts and may also be found in muscle after considerable activity, most of it does not exist as such in perfectly relaxed muscle, but occurs here as phosphocreatine frequently about 2 millimoles per 100 g. The discovery of

phosphocreature was preceded by the observation that part of the apparent inorganic phosphate in muscle was in reality present as a labile compound, called phosphagen, which was hydrolyzed by molyhdic acid under the conditions of the phosphate determination. This phosphagen was then identified as phosphocreatine by Fiske and SuhbaRow. In in-



FIO 73 CREATINE

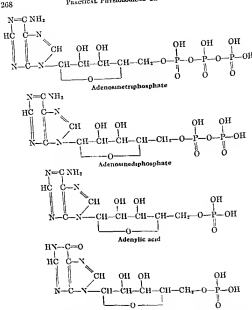
vertehrate animals, other phosphagens occur, among these, phosphoarginine has been shown to be of widespread occurrence

Purme Bases and Nucleotides. The purme hases occurring in muscle are adenine, guanne, vanthine, and hypovanthine (for formulas see Chapter 7) These may exist in the form of nucleic acids and of mono- and obgonucleotides and their hreakdown products. Among these, we shall discuss only the adenosine polyphosphates.

Iu all cases, the hasic structure is adenosine, adenine-9-rihoside, n(-)rihose being the carhohydrate moiety. This is esterified with phosphoric
acid at the 5 position of the rihofuranose, to form adenosine-5-phosphoric
acid, also known as adenosinemonophosphate (AMP) or as muscle- or
myoadenylic acid. Inosinie acid is a commonly occurring hreadown product of AMP, formed by deamination in muscle extract. Though the nature of adenylic deaminase has not been exhaustively investigated, it
appears that myosin, even after considerable purification, displays such
enzymic activity.

By attaching further phosphoric acid residues in pyrophosphate linkage adenosinediphosphate (ADP) and adenosinetriphosphate (ATP) are obtained Iu resting muscle as well as in the minimalian heart, ATP is

¹ Kanner Fortschr Chem. org \aturstoffe 8, 96 (1951)



Inosinic acid the predominant form of the adenosine polyphosphates usually in a con-

centration of about 0 5 millimole per 100 g Carnosine. This substance is a dipeptide of histidine and the amino acul β-alanine, which occurs also in pantothenic acid (see p 35) Carno sine, often accompanied by anserine (methylcarnosine), is of widespread

Carnosine

ocurrence in vertebrate muscle, but its function is not known 2 ² du Vigneaud and Lehrens Ergebn Physiol 41, 917 (1939)

Carnitine is a betaine which gives trimethylamine on hydrolysis. Its

function in muscle is not known

Glutamine. Skeletal and, particularly, cardiac muscle contain this substance in higher concentration than other mammalian tissues,3 about 0.1 per cent. Its function is not known, but reference may be made to the discovery by Krehst that glutamine is essential for the maintenance of the intracellular potassium in brain (For formula of glutamine, see p. 145)

INORGANIC SALTS

The predominating inorganic cation of muscle is potassium, occurring in a concentration of about 0 11 molar It is accompanied by sodium, which, like the chloride ion, is mostly present in the tissue spaces rather than jutracellularly. Also significant are calcium and magnesium (about 0 003 and 0 01 molar), which act as activators or inhibitors for many of the enzymes present These positive ions act in part as counterious for the negatively charged proteins, and are, for the remainder, paired with various phosphoric esters and other amons. Free phosphate occurs, but is easily overestimated due to the lability of phosphocreatine. The actual estimates differ for different muscles, but the results of Hastings for rat muscle may be taken as representative for mammalian muscle is

APPROXIMATE	COMPOSITION	OP 1	-0	On 3	1	Margara

1	C	Villiequivalents		Distribution, per cent		
	Grams	Cations	Antons	Extracellular	Intracellular	
Water Protein	760 220		~50	20 10	80 90	
Na K Ca Mg Cl HCO ₁ Phosphates		21 107 3 22	12 10 ~80	80 1 >90 1 >99 40	20 99 <10 99 <1 60 >99	
Total		153	~152			

PROTEINS OF MUSCLE

The protein composition of muscle must be discussed in relation to the microscopic structure of this tissue. We can divide the proteins into those

Archibald Chem. Rev. 37, 162 (1945)
 Terner Eggleston, and Liebs Biochem J. 37, 139 (1950) Hastings, in Najiar A Symposium on the Chinical and Biochemical Aspects of Carbo-hydrate Utilization in Health and Disease Baltimore, The Johns Hopkins Press 1959

that are constituents of the fibrils and those that occur in the sareoplasm, even if in practice it may not always be possible to make this distinction with certainty

Stroma. About 20 per cent of the muscle protein cannot he dissolved with any known solvent short of strong urea or alkali solution. This fraction is commonly called stroma, and may consist, in part, of constituents of cell nuclei and connective tissue, including perhaps the sarcolemnas sheaths.

Proteins of the Sarcoplasm. If a muscle is perfused to remove blood, and is then mineed and subjected to high pressure, a press juice is obtained which contains about one-fifth of the protein of the mineed muscle. This press juice is probably derived from the sarcoplasm. Its main proteins were formerly called myogen and globulin X (Weber), the latter protein has not received further attention. *Upogen* is now recognized as a collective name for a number of water-soluble proteins, and it is probable that this fraction is made up entirely of various enzymes.

In more complete yield, the proteins of the myogen group can be obtained by extracting mineed muscle with water or dilute salt solutions (e.g., 0.15 molar KCl). It is then found that about 30 per cent of the muscle protein is contained in this fraction. Such extracts have been analyzed with the ad of electrophoresis, and this method too has shown the com-

pleaty of the so-called myogen (Dubusson)
Myoglohin. A water-soluble protein which deserves separate mention
is myoglobin. This protein is similar to hemoglobin (notwithstanding differences in molecular weight (17,000) and other properties) in that it is a
heme protein which combines reversibly with oxygen. Its affinity for Os
is higher than that of hemoglohin, so that the intracellular pigment may
be fully oxygenated at O₃ tensions which cause unloading of the blood
pigment. The distribution of myoglohin between white and red muscle
bas most interesting features correlated with physiological differences between the muscles. It may be said, in general, that myoglobin acts as a
store of oxygen, although the functional meaning of this is not always
clear.

Myoglobin can be released from muscles after crushing injury, because of the small size of the molecule, it is filterable through the renal glomeruli, and appears in the urine

Proteins of the Fibrils. The two previous categories constitute nearly half of the muscle protein, the remaining half will now be discussed in detail. This fraction, which corresponds to the protein constituents of the fibrils, can be extracted with concentrated salt solutions (e.g., $0.5~\mathrm{M}$ KCl), and bas been extensively investigated

Myosin. When relaxed skeletal muscle is minced and immediately extracted with 0.5 M KCl for an arbitrary length of time at neutral or weakly alkaline reaction it is found that the extract, in addition to the soluble sarcoplasm proteins contains a protein, called myosin, which can be precipitated by five fold to ten fold dilution of the extract, and can be reduscolved in 0.5 M KCl Myosin solutions thus obtained are distur-

⁴ Herbert Gorl n Subrahmanyan and Green Buchem J 34, 1108 (1940)

guished by a high viscosity and flow birefringence,7 but these properties are found to be variable and are largely determined by the duration of the extraction, prolonged extraction gives rise to a myosin with higher viscosity and more intense birefringence of flow 5

Actin. Further analysis has shown that these phenomena are due to the combination of myosin with a new protein, actin, to form the compley, actomyosin This combination takes place in later stages of the extraction as soon as the adenosmetriphosphate in the system is decomposed, but the release of actin is also promoted by very fine mincing Pure, actin-free myosin has been prepared in crystalline form by Szent-Gyorgyi Its viscosity is moderate, and it shows no flow hirefringence when tested with simple methods of observation. Upon combination with actin, however, the viscosity increases greatly, and flow hirefringence is pronounced

Actin can occur in two forms, called globular (G) and fibrous (F) actin The former can be obtained by extracting acctone-dued muscle nowder with water After addition of salt, the G-F transformation takes place, as evidenced by an enormous increase in viscosity and the development of a strong flow birefringence * These phenomena are ascribed to a suppression of molecular charges by the salt, whereupon the individual molecules unite to form long molecular strands 10 Only I-actiu forms an actomyosin with the described characteristics

Both actin and myosin are now available as apparently pure proteins For myosiu, purity is achieved by carefully controlled precipitation procodures The molecular weight of my osin has been difficult to determine. owing to anomalies in its behavior, but is now found to be 600,000, the molecule apparently heing a rod 1500 Å long 11 Actin has been purified hy fractional ultracentrifugation of F-actin followed by careful depolymerization to G-actin, the molecular weight is about 57,000 1°

Much remains to be elucidated about the exact state in which these proteins occur in the muscle fibril Electron microscopy has shown, however, that long filaments, presumably actomyosin, run along the fibrils, and that other substances are spaced in a regular fashion, corresponding to the microscopically visible cross strictions and other structural details 13

Tropomyosin. This protein was crystallized by Bailey after extraction from alcohol dried muscle. Its behavior is opposite to that of actin in that it is aggregated in the absence of salt, but has its flow birefringence abolished by salt However, it is not known to be related to contractile

There is a good deal of similarity in amino acid composition between my osin, tropomyosin, and actin Apparently, these proteins are built on another pattern than seed, e.g., or blood proteins

on Muralt and Edsall J Biol Clem 89 315 351 (1935)
Banga and Szent Gyorgyi Stud Inst Wed Chem Seged 1 3 (1942) Straub Stud Inst Vici Chem Szeged 2 3 (1942) 3, 23 (1943)

¹⁶ Mommaerts J Biol Chem 198 459 (1952)
¹¹ Mommaerts and Rupp J Biol Chem (in press)

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BIOCHEMISTRY OF MUSCULAR ACTIVITY

Our present knowledge of the chemistry of muscular contraction is based upon the intensive studies of many investigators over a long period of time. Perhaps no other single phase of animal biochemistry has gogests Muscles from both warm- and cold-hlooded animals have been studied; skeletal, cardiac, and (to a lesser extent) smooth muscles have been used; and studies have heen carried out on the isolated intact muscle, muscle extracts, and enzyme systems obtainable from these extracts. The discussion which follows is a summary of present knowledge, necessarily incomplete and subject to future revision, concerning the chemical processes associated with muscle contraction.

CLYCOLYSIS

Glycolysis is the preponderant metaholic process in muscle under anaerolus conditions, but may also occur aerobically in those muscles which do not contain enough oxidative enzymes to fulfill the metaholic requirements of increased activity. In the human hody, considerable amounts of lactic acid appear in strenuous exercise; much of this lactic acid is reconerted to glycozen by the liver.

Glycogenolysis. Breakdown of glycogen can be effected hydrolytically by amylase, but this occurs in digestion rather than in tissue metabolism. In muscle and other organs, the hreakdown is phosphorolytic instead of hydrolytic, and is catalyzed by the enzy mo phosphorylase, which has been

isolated in crystalline form14 and studied extensively in Cori's laboratory.

This reaction may be visualized as shown on p. 272.

This reaction has a number of interesting features. It is an equilibrium reaction which leads to synthesis of glycogen if there is sufficient glucose phosphate relative to phosphate, whereas breakdown occurs in the opposite case. The synthetic reaction requires a small amount of glucose polymer as a primer. In this phosphorylase reaction, an unbranched chain polymer is formed, with all glucose molecules linked by 1.4 linkages; this

product is a linear starch, which stains blue with iodine. The synthesis of branched polymers with both 1,4 and 1,6 linkages (of which glycogen is an extreme example) requires the additional participation of a branching

factor. This enzyme causes the transfer of parts of straight chains to nonterminal positions. Phosphorylase caunot break 1,6 linkages; these are

hydrolyzed by a special enzyme.

Phosphorylase is known in two forms, a and b. The first of these is active as such; but phosphorylase b requires the addition of adenylic acid, which seemingly acts as a loosely combined cenzyme. It is probable that phosphorylase a, too, contains a nucleotide as its active group, but this is bound strongly, and is not identical with adenylic acid. Various tissues contain an enzyme that converts phosphorylase a into b; this conversion is observed in muscles after exhaustive activity.

Glucose-1-phosphate is transformed into glucose-6-phosphate by the enzyme phosphoglucomutase, crystallized by Najjar; glucose-1,6-diphos-

¹⁴ Green, Corn, and Cori; J. Biol, Chem , 142, 447 (1942).

phate acts as coenzyme for this reaction. The blood glucose is also a source of glucose 6-phosphate in muscle The transformation cannot be effected by direct combination of glucose with phosphate, since the equi librium does not favor synthesis Instead, glucose is phosphorylated by ATP under the influence of the enzyme hexol inase

Formation of Triosephosphate. Glucose-6-phosphate, whether formed from gly cogen or glucose, is transformed into fructose-6-phosphate in the presence of the enzyme oxoisomerase, and the reaction product is phosphorylated once more by ATP to form fructose-1,6-diphosphate These steps are shown on p 273

Fructose-1,6-diphosphate is then split under the influence of the enzy me aldolase into two isomeric triosephosphates, dihydroxy acetonephosphate

and 3-phosphogly ceraldchy de

These triosephosphates are interconvertible through the action of an other isomerizing enzyme It is the 3 phosphoglyceraldehyde which under-

goes further reaction

Oxidative Step. Triosephosphate is oxidized by pyridine nucleotide (DP\, coenzyme I) and the enzyme phosphoglyceraldehyde dehydrogenase, in a reaction which would be expected to yield 3-phosphoglyceric acid Instead, it is found that 1 3-diphosphoglycerate is formed, while morganic phosphate (without which the reaction cannot proceed) is taken up (War burg) One may consider that the dehydrogenation of aldehyde, normally my olving a preliminary hydration, in the present case utilizes phosphoric acid instead of water to that effect. The actual reaction mechanism is a problem of considerable interest 14

Formation of Pyruvate. Diphosphoglycene acid is transformed into 3-phosphoglyceric acid, ADP acting as the phosphate acceptor The product is converted into 2-phosphoglycerate and then, by the enzyme enclase, into enol-phosphopyruvie acid This finally donates its phosphate radical to ADP, yielding pyruvate COOH

Formation of Lactate. In the formation of diphosphoglycerate from triosephosphate, a molecule of coenzyme I had been reduced. This is now reondized by pyruvate, reducing the latter to lectic acid, which is the end product of glycolysis CH₃

RESPIRATION

Under aerohic conditions, essentially the same set of reactions takes place as in glycolysis, up to the stage of pyruvate formation However, the reoxidation of coenzyme I reduced in the triosephosphate dehydrogenation is undertaken by oudative enzymes and eventually by oxygen Pyruvate then is not reduced to lactate Instead, it is completely oxidized

in the citric acid cycle (see Chapter 33)

The respiratory euzymes, like cytochromes and cytochrome oxidasc (see Chapter 12) are present in different muscles in varying amounts, although muscular activity eventually depends upon respiratory metabohsm There are types of muscle, like the myocardium, is which are coutinuously active and must therefore steadily respire at a sufficiently high rate to meet the energy demands Such muscles must contain sufficient respiratory enzymes to permit this high rate of metabolism Other types of muscle, whose activity alternates with periods of rest, frequently are less well endowed with oxidative systems. When such muscles become

¹⁴ Much of the classical work on cytochromes, cytochrome oxidase yellow enzymes and other oxidation factors was done on preparations of mammalian and pigeon heart muscle

active, glycolysis takes place in addition to respiration, in an effort to fulfill the energy requirements

ENERGY RICH PHOSPHATES

Neither oxygen nor lactic acid is directly involved in contractility since muscles can contract anaerobically when glycolysis is inhibited with indoacctate. Furthermore, increased oxygen consumption and lactic acid production occur only after contraction (Embden). Phosphocreatine and ATP are found to be more closely related to the contractile mechanism. These compounds are examples of the so-called high-energy phosphates, compounds which liberate energy when dephosphorylated, as manifested by the large amount of heat produced in their hydrolysis (about 12 000 calones per mole hydrolyzed). This applies to phosphocreatine and to each of the two pyrophosphate linkages in ATP, but not to the rhose-phosphate linkage. The free-energy effect of the sphtting is likewise highs that such compounds can act as phosphate-group donors for the synthesis of other substances, and can presumably, also serve to yield energy to be transformed into mechanical work.

Generation and Utilization of ATP. The description of the reaction sequence of glycolysis has given examples of how, in the formation of glucose-6-phosphate from glucose and of fructose-1.6-diphosphate from fructose-6-phosphate, ATP acts as a phosphate group donor, and how in other instances reaction products like diphosphoglycerate or phosphopyruvate transfer phosphate groups to ADP It is indeed, the purpose of metabolism to subject the metabolites to such transformations that the phosphate groups contained in them acquire a high group potential so that they can be transferred to ADP (or AMP, although this, in general, seems to be less important) By such reactions, ATP is regenerated when ever it has been split in other reactions. In glycolysis, 4 moles of ATP are thus formed per mole of hexose decomposed, 2 moles having initially heen expended to phosphorylate glucose to the hexose diphosphate stage (only I mole in the case of Llycogen breakdown but the formation of glycogen from blood sugar also expends I mole of VTP) In respiratory metabolism there are many other reactions in which VTP is produced not all of which are known in detail Aerobic phosphorylation may lead to about 6 moles of ATP generated per mole of Oz consumed

Enzymes and ATP. A number of enzymes are known which are of in trust in connection with the transformations of VTP. The following enumeration is not complete but is restricted to some enzymes occurring in muscle. Hydrolysis of ATP is effected by adenosinetriphosphatases or MTP asses which split the terminal pyrophosphate linkage, yielding in organic phosphate and VDP. W. least two such craymes occur in muscle one of these is my osin itself (Engelhardt and I jubimova), the other is a particulate entity similar to the microsomal or mitochondrial fractions isolated from other organs (Kielley and Weyerhof). Both enzymes display a compleated dependence on the ion composition of the medium. The particulate VTP ase is activated by magnessum and inhibited by call.

Lap narn Adrances in Enzymol 1 99 (1911)

cium, hut for pure myosin the opposite is true. In actomyosin systems the situation is more complicated and has not yet heen fully explored. The function of the particulate enzyme is probably not related to contractility. Myosin, on the other hand, is a part of the contractile machinery, hence its ATPase activity is a matter of obvious interest. However, it has been stated that the activity of myosin is not sufficient to account for the actual rate of breakdown occurring in sustained tetanic activity. It is probable that the ATPase activity of myosin is but a reflection of a more fundamental reaction which has not yet been recognized, and which may be related to the occurrence of myosin in an organized state in conjunction with other active materials.

Phosphocreatine, on the other hand, is not directly hydrolyzed by musele enzymes. It can transfer its phosphate group to ADP (and also to adenvise acid, AMP) in the Lohmann reaction.

so that dephosphorylation of phosphocreatine can be effected by a system containing creatine-ATP-phosphopherase ATPase, and adenine nu eleotide It is held that in living muscle phosphocreatine acts as a reservoir of ~ph, from which ATP can be restored on short notice A more fundamental role of phosphocreatine is however, not at all excluded, and it may be specifically involved in relavation (Duhusson)

ATPase removes only one phosphate from ATP, but an enzyme exists (myokinase) which catalyzes the transmutation 2 ADP = ATP + AMP In combination with ATPase, this enzyme can thus effect the complete dephosphorylation of ATP to AMP, which, as mentioned previously, can then he deaminated to mosine and (IMP). The latter is formed after prolonged muscular activity, but it is doubtful whether such far going breakdown occurs normally under optimal conditions. Formation of IMP may be the normal pathway of autolytic nucleotide catabolism in skeletal muscle, but not in heart or other tissues where adenytic acid is dephosphorylated rather than deaminated (Kerr).

Other enzymes effect the transfer of phosphate from ATP onto other substrates, thus, hexokinase catalyzes the reaction ATP + glucose \rightarrow ADP + glucose \rightarrow ADP + glucose \rightarrow ADP and the possible connection of such enzymes, instead of ATPases, with activity metabolism has not yet received sufficient consideration.

True and Apparent ATPase Activity. When an isolated enzyme, myosin or some other, splits ATP mto ADP and phosphoric and, without demonstrable participation of other substances, it is no doubt correct to call such an enzyme "adenosinetriphosphatase" It is an interesting question whether this nivolves phosphorylation and dephosphorylation of the enzyme protein as intermediate steps, but this would not affect the terminology. It is also possible that in more complex systems a set of reactions would occur as follows.

$$ATP + X \rightarrow ADP + X-ph$$

 $X ph \rightarrow X + phosphate$

In such a case, X would be a coentyme of the VIP ise.

It should be understood, bowever, that if a splitting of ATP is observed in a crude tissue extract or in an intact organ, this splitting is not necessarily due to an ATPase, with or without coenzyme, but may be linked with other and irreversible metabolic reactions. Thus, the formation of urea in liver from proper ammonia donors through the functioning of the ornithine-citrulline-arginine cycle (see Chapter 33) involves the splitting of ATP Likewise, formation of glucose-0-phosphate from ATP and glucose, followed by breakdown of the glucose phosphate or its conversion products, would appear as a hydrolysis of ATP Such additional metabolic components have been referred to as cosubstrates. In The breakdown of ATP in complex tissue systems may well be due to such reactions rather than to ATPase.

ATP Breakdown and Muscular Activity. It was shown by Lundsgaard that, when respiration and glycolysis are chminated, muscular activity is linked with the dephosphorylation of phosphocreatine As explained, this proceeds presumably via ATP breakdown. The primary breakdown of ATP was demonstrated recently by comparative analyses of relaxed and contracted muscle, both striated and cardiac 19 It appears that in each case one cyclo of activity is linked with the decomposition of maximally 5 × 10-7 moles of ATP per gram of muscle, as predicted on the basis of physiological data. Thus, we regard the dephosphorylation of ATP as the metabolic reaction primarily connected with contraction The study of heat production in a single twitch (Hill) supports this sams sequence the contraction heat corresponds to splitting of ATP, relaxation (restoring ATP from phosphocreatine) is almost thermoneutral, and the restitution beat depends on the provalent type of metabolism, whether glycolytic or respiratory In the more physiological case of tetanic contraction, the various phases overlap

Although ATP breakdown has been thus correlated with contraction, nothing warrants the assumption that this breakdown is to be ascribed to direct ATPase activity, rather, it may involve complicated cosubstrate

reactions or mechanochemical mechanisms

MOLECULAR MECHANISMS

In muscular contraction, chemical energy is directly transformed into mechanical work, and this transformation takes place in the muscle fibril. An explanation for this phenomenon could be given if it were known just what the molecular structure of the fibril is, how this structure is modified to result in contraction and how this modification is connected with the breakdown of ATP. This objective has not been reached, much less the corresponding analysis of relaxation and of neuromuscular activation. A number of interesting advances, however, warrant the hope that

an evolunation of contraction in terms of actomy osin-ATP interaction may eventually be achieved

It is possible to prepare threads of actomy osiu by extrusion of a solution of this protein into a dilute salt solution Such threads contract when ATP is added, especially in the presence of some magnesium 20 Oriented and stronger threads can be made by various devices, these thicken durmy contraction and can lift weights Finally, one can prepare hundles of actomyosin filaments in their original configuration by extracting all other constituents from muscle strips Such fiber preparations become thicker when shortening and develop about the same tension as the original muscle would *1

Imperfect though the extruded actomy own fiber is, it permits the study of the individual protein components. Thus, it has been found that neither myosin nor actin alone can form contractile fibers, but that only the combination of myosin with F-actin is capable of contraction Apart from the cation reonirements, it can be said, then, that contractility is the result of the effect of ATP upon an oriented structure of actiu and myosin. The mechanism of this unteraction is unknown, but the formulation of the problem in these biochemical terms is itself a great advance

In addition to its role in combination with myosin, actin itself displays some interesting properties. The polymerization of actiu from the G to the F state is linked with the breakdown of ATP in a stoichiometrio reaction

The amount of actin in one gram of muscle is such that, if it should all polymerize at one time, 5 × 10 1 moles of ATP would be split, which is exactly equal to the amount of ATP broken down in one maximal contraction *2 This equality makes it likely that polymerization takes place early in the activation phase of the contraction, to he followed by contraction of the F-actomyosu formed

The participation of actomyosin and its components in contraction is further illustrated by the observation that these proteins can be extracted only from relaxed muscle Contracted muscle yields no actomyosin or myosin, instead there appears a new protein, contractin, albeit in smaller amount (Dubuisson)

EXPERIMENTS ON MUSCULAR TISSUE

SEPARATION OF EXTRACTIVES FROM MUSCLE

1. Creatine Dissolve about 10 g of a commercial extract of meat" in 200 ml of warm water (Test for protein by biuret and congulation tests, pp 171, 188) Precipitate the inorganic constituents by neutral lead acetate, being careful not to add an excess of the reagent (30 ml of a 20 per cent solution is about the right amount) Write the equations for the reactions taking place

^{**} Szent-G) orgy: Stud Inst Med Chem. S.eped 1 17 (1942)

**Szent-G) orgy: Biol Bull. 96, 140 (1949)

*** Mommaerts J Biol Chem. 198, 469 (1952)

¹¹ Commercial meat extracts vary considerably in their creatine content and may have to be fortified for class use.

here Allow the precipitate to settle, then filter, and remove the excess of lead in the warm filtrate by hydrogen sulfide Fliter while the solution is jet warm, evaporate the clear filtrate to a syrup, and allow it to stand at least 48 hours in a cool place. Crystals of creatine should form at this point Examine under the microscope (see Fig. 73) Treat the syrup with 25 ml. of 95 per cent ethyl alcohol, stir well with a glass rod to bring all soluble material into solution, and then filter. When the fluid has drained completely, use 2 further 10-ml portion of 95 per cent alcohol to ald in transferring the residue In the beaker to the filter paper, and tn wash the material on the filter paper. The purine bases have been dissolved and are in the filtrate, whereas the creatine crystals were insoluble in the alcohol and remain on the filter paper. Remove the crystals and bring them into solution in about 10 ml. of bot water. A little animal charcoal may be added to decolorize the solution, Filter and concentrate the filtrate to small volume. Allow the solution to cool and note the separation of coloriess crystals of creatioe.24

Make the following tests on the crystals.

a Microscopical Examination Examine some crystals under the microscope and compare the form with those reproduced in Fig. 73

b TRANSFORMATION OF CREATINE INTO CREATININE. Dissolve a portion of the crystals in about 10 ml. of water. Divide into two equal portions. To one portion add 5 ml of 2 N hydrochloric acid. Evaporate this acidified portion carefully over the free fiame and finally to dryness on the water hath. The creatine has been chaoged into creatinine. Take up the residue in about 5 ml. of hot water, cool, and apply the tests for creatinioe as given in Chapter 28 to this extract as well as to the original solution. What are your conclusions?

Diacette Reaction To 5 ml, of a dilute creatine solution add an equal volume of saturated sodium carbonate solution and a few drops of a solution of diacetyl A pink color should develop. This test has been made the basis of a method for the quantitative determination of creatine (see p 287).

2 Preparation of Glycogen " Grind a few fresh oysters in a mortar with sand 14 Transfer to an evaporating dish, add water, and boil for 20 minutes. At this point the volume of solution should have been reduced by about onehalf Note the opalescence of the solution At the bolling point, faintly acidify with acetic acid. Why is this acid added? Filter, and divide the filtrate into two parts

Test one part of the filtrate as follows.

a loding Test To 5 ml. of the solution in a test tube add 5 to 10 drops of more of Lugoi's sodine solution, at the same time adding a similar amount of iodine to 5 ml of water in another tube, this serving as a control. What do you observe? Is this similar to the lodine test upon any other compound with which we have had to deal?

b Reduction Tests. Does the solution reduce Benedict's solution?

e Hypholysis of Glycogen Add 10 drops of concentrated hydrochloric acid to 10 ml of the solution and boil for 10 minutes Cool the solution, neutralize with solid sodium carbonate, and test with Benedict's solution. Does it still

^{**} For the preparation of pure creatine from creatinine see p 801

is For a quantitative experiment showing the effect of diet on the glycogen content of hver see Chapter 33 p 1071

[&]quot;Clycogen may be readily obtained from the livers of well fed rabbits which have been killed by the intraperitoneal injection of 5 ml of 25 per cent anhydrous MgSO4 per ke The excised liver is quickly cut into small pieces and dropped into about 150 ml of boiling

fail to reduce Benedict's solution? If you find a reduction how can you prove the identity of the reducing substance?

d. INFLUENCE OF SALIVA Place 5 ml. of the solution in a test tune, add 5 drops of saliva, and place in the water bath at 40° C. for 10 minutes. Does this now reduce Benedict's solution?

To the second part of the glycogen filtrate add 3 to 4 volumes of 95 per cent alcohol. Allow the glycogen precipitate to settle, decant the supernatant fluid, and filter the remainder. Transfer the glycogen to a watchglass and heat on a water hath to remove the alcohol; then subject it to the following tests:

- a. Solubility. Try its solubility in cold and hot water, in alcohol, and ether.
- b. IDDINE TEST. Place a small amount of the glycogen in a depression of a test tailet and add 2 to 3 drops of dilute iodine solution. The same wine-red color is observed as in the foldine test upon the glycogen solution.

salt has the empirical formula G_1 II₁ $O_1N_1P_2Ba_2 + xII_1O$, the water content depending upon the extent of drying Purity is usually established by the following criteria (1) The molecular ratio of nitrogen to phosphorus, as established by total N and total P analyses, should be 5 3, (2) the inorganic phosphate content should be very low, (3) the "labile phosphate" (see below) should be two thirds of the total phosphate after correcting the latter for any long and phosphate present

- 2 Conversion of Barium ATP to Sodium ATP. Dissolve 200 mg of the bar lum sait in 6 ml of lee cold 0 1 N hydrochloric acid in a centrifuge tube 460 mg of anhydrous sodium sulfate dissolved in a little water, sitr, and can trifuge. Decant the supernatant and wash the precipitated barium sulfate several times with small portions of cold water containing a trace of hydrochloric acid. Neutralize the combined supernatant and washings to pil 7.5 8 5 as desired, analyze an aliquot for inorganac and labile phosphate, and as the basis of this analysis didute to the desired concentration of sodium ATP. This solution is relatively stable for some weeks in the cold, but undergot slow hydrolysis, the extent of which should be established before use
 - 3 Conversion of Ba-ATP into Soluble Solis by Vecons of a Cation exchange Ren Dissolve 200 mg of the barlum salt in 6 ml cold 0 l N hydrochord eard, and dilute to about one half of the volume eventually desired Prepart a column of 3 cm height, in a tube of about 1 cm width (closed near the bottom with a plug of glass wool or a firsted glass disc), of a cation exchange real in the potassium or sodium cycle 2 Filter the solution through the column at a rate of a few mil per minute, and wash the column with water The emerging solution is devoid of barum but it susually still slightly and it is adjusted to the desired pil with KOH or NaOH and brought to the required volume

This procedure is more convenient than the precipitation of banum si sulfate, and is also effective in removing traces of heavy metals which would interfere in many enzyme experiments ? content, corrected for the inorganic phosphate already present, represents the "lahile phosphate" of ATP. It should be two-thirds of the total ATP phosphorus. 2 (3) Total puosphorus. Transfer a 1 ml. portlon of the ATP solution to a large test tube or micro-Kjeldahl flask and add 2.5 ml, of 5 N sulfuric acid. Heat over a microburner until the water has boiled off and the residual fluid is brown or black. Add 1 drop of 30 per cent hydrogen peroxide and holl until clear. Cool, transfer to a 25-ml, graduated flask, with washings to about 18 ml. volume. Add 2.5 ml. of ammonlum molybdate and 1 ml. of aminonaphtholsulfonic acid reagent. Dilute to 25 ml., mix, and allow to stand 10 minutes. Compare in a photometer against a standard containing 0.1 mg. P digested with acid, etc., just as was the unknown. A blank of water alone should also be run through the entire procedure, to correct for any phosphorus in the reagents. The total P of the sample, corrected for any Inorganic P present, represents ATP phosphorus. (4) Total MINROGEN, Determine by any suitable micro-Kieldahl method (see Chapters 23 and 31). A 4-ml. portion of the ATP solution described above will contain about 0,3 mg, of nitrogen. The N:P ratio on a molecular hasis should be 5:3. (5) ULTRA-VIOLET ABSORPTION. Measure the absorption spectrum of a suitably diluted neutral or acld solution of ATP (containing about 10 to 40 mg, per liter) hetween 230 and 300 mg. The absorption maximum should be situated at 258 to 260 mg. The adenine content can be calculated on the basis of an extinction coefficient of 1.6 at the maximum for a 10-4 molar solution. The adenine:N:P ratio should be 1:5:3, on a molecular or atomic basis.

5. Preporation of Adenylic Acid from Adenosinetriphosphote (Kerr), Dissolve 2.5 millimoles of barlum adenosinetriphosphate (93 mg. of ATP phosphorus equals one millimole) in a flask containing about 100 ml. of 0.1 N hydrochloric acid, and add a few drops of phenolphthalein followed hy sufficient barlum hydroxide solution to produce a permanent pink color. Attach a reflux condenser to the flask and beat to holling over a free flame. Boil for 30 minutes, maintaining the reaction just aikaline to phenolphthalein by the continuous addition of barlum hydroxide solution in small portions through the condenser tube. Cool to room temperature, add sufficient N bydrochloric acid to the flask contents to dissolve the precipitate, and dilute to about 1,750 ml. with water. Again neutralize to phenolphthalein with harium hydroxide, and allow the precipitate to settle. Remove the supernatant by decantation and filtration, discarding the precipitate. To the supernatant add sufficient acetic acid to make the final acid concentration 0.2 per cent, followed by 50 ml. of 20 per cent mercuric acetate solution per liter of fluid present. Allow the precipitate to settle overnight, separate it by decantation and centrifugation, and wash it once with 0.5 per cent mercurio acetate solution. Suspend the washed precipitate in about 50 ml. of water containing a few drops of 2 N sulfuric acid and treat with hydrogen sulfide for one hour. Filter off the precipitated mercuric sulfide and pass air through the filtrate to remove excess hydrogen sulfide. The volume at this point should not be over 25 ml, per millimole of starting material. Add acctone to the solution at 20° C, to the first permanent turbidity (not over 1.5 volumes). Allow to stand on ice overnight, filter and discard the precipitate. Bring the filtrate to room temperature and again add acetone to the point of turbidity. Again chill overnight. Separate the crystalline adenylic acld by centrifugation. Repeat the addition of acetone as described until a total of 3 volumes has been

¹¹ Determination of the acid-lable P (2 moles per mole ATP) is a method frequently used for the determination of ATP. It is a popular misconception that this can also be used to prepare adea; in acid. In reality, the nucleotide is split by this procedure into adenine, ribose-5-phosphate, and morganic phosphate.

added Drive off excess acetone fram the separated adenylic acid by gentle warming. Recrystallize by dissolving in the inhimial amount of het water filtering quickly, and allowing the filtrate to stand in the cold overnight. Centrifuge or filter off the crystals and repeat the recrystallization procedure five or air more times to obtain a final product which has the theoretical N P ratio of 5 1 and the correct melting point of 189° C. The final crystals are washed with a little alcohol and other and dried. The mother liquors from the various recrystallizations may be assed, combined, and treated with 3 volumes of acetone, followed by recrystallization as above, to obtain more adenyie acid which is less pure

PREPARATION AND REACTIONS OF MYOSIN AND ACTIN

- 1 Preparation of a Crude Myosin Solution Anesthetize and kill a lean rabbit, as described for the preparation of ATP, and chill and excise the musculature in the same manner Miner the tissue in a chilled meat grioder and extract immediately for 20 minutes in the cold with 3 volumes of 0.6 M KCl 0.05 M NaHCO, Collect the extract by centrifugation or straiologisthrough several layers of gauze and dilute with 10 to 12 volumes of cold water (redistined in a glass still or purified through a mixed bed ion exchange resin, in a so called water demineralizer which should be made of plastic without metal). Allow the precipitated myosin to settle in the cold siphon off most of the supermatant solution, and collect the myosin by centrifugation Redissolve the myosin by the addition of a measured amount of 2 M KCl concentration is 0.5 M
 - 2 Formation of Actomyouth Proceed with the extraction of minced muscle as in the previous experiment, but incubate the tissue with the alkaline KCI solution for various lengths of time up to 18 to 24 bours in the cold, with occasional stirring Take 50 ml aliquots at regular intervals, and obtain the extracts from these by centrifugation followed by fittration through glass wool Formation of actomyosin is indicated by a strong rise in viscosity and flow birefringence (see below) Actomyosin can be prepared from the extract by precipitation and resolution as described for myosin
 - 3 Observation of Flow Birefringence: A simple apparatus can be assembled with the aid of two 2 incb polaroid discs mounted vertically above each other about 4 to 6 inches apart and illuminated from below. The polaroids are crossed, so that no light is transmitted A 30 ml glass heaker is selected such that when examined in the polaroscope thus assembled, its bottom does not show significant birefringence. A crude or purified solution of actomyoun is placed in the beaker to about 1 cm depth and this solution, while being observed between the crossed polaroids, is gently rotated by swifling the beaker by band. The rotating solution shows a dark cross (the isocilyne cross) on an Illuminated background. This phenomenon is due to the orientation of very long particles (molecules or swarms) in the streaming gradient which leads to a regular alignment giving rise to hirefringence, except for those orientations that are parallel with the vibrational axes of the polarizer and
 - 4 Preparation of Crystalline Myosin 11 Prepare minced rabbit muscle 28 previously described and extract 309 g in the cold with 1 liter of 0.3 M KCl

²³ This procedure comianes different features of the methods of Szent-Gyorgyi of Weber and of Mommaerts and Larrah.

0.15 M K-phosphates (pH 6.5) for 15 minutes with gentle stirring. Add 4 liters of cold water, and strain through several layers of gauze. Dilute with water to 12 liters final volume, and leave in the cold for 3 hours. Siphon off the supernatant solution as far as possible, collect the protein by centrifugation, dissoive it by the addition of 60 ml. of 2 M KCl and 30 ml. of a solution 0.25 M with respect to K-HPO, and KH-PO, each, and add water to a final volume of 300 ml, (pH 6.5 to 6.8, ionic strength 0.5). Clarify this solution by centrifugation and filtration through glass wool or Filter-cel. Add 240 ml. of water, and remove the precipitated actomyosin by strong centrifugation. (The solution may remain somewhat turbld through micellar aggregation of myosin at this ionic composition.) Place the solution in a large, cooled container and slowly (10 minutes) add 4 liters of cold water with continuous stirring. The myosin will precipitate in the form of fine, microscopically visible needles which give a silky sheen to the suspension. The steps for the removal of actomyosin and the crystallization may be repeated, and the myosin finally dissolved in 0.5 M KCl, and dialyzed against this solvent.

Pure myosin shows no flow hirefringence with the simple method described in Experiment 3 of this section.

- 5. Demonstration of Adenosinetriphosphatase Activity of Myosin Preparations. Place 2 ml, of myosin solution (diluted in 0.5 M potassium chloride so as to contain 1 to 2 mg, of protein) in each of two test tubes. Add 1 ml. of 4 per cent sodium bicarbooate solution and 0.5 ml. of 0.1 per cent anhydrous calcium chloride solution. To Tube 1, which is n control, add 0.5 ml. of 20 per cent trichloroacetic acid solution. To each tube add 0.5 ml. of sodium ATP solution containing a known amount (0.1 to 0.2 mg.) of "labile phosphorus." Dilute the contents of the control tube immediately to 10 ml. with water, mix, and filter. Allow the second tube to stand at room temperature for 30 minutes, then add 0.5 ml. of 20 per cent trichloroacetic acid solution, dilute to 10 ml., mix, and filter. Determine the Inorganic phosphate content of each filtrate as described above, using a 2 ml. aliquot. Correct for the value of the control by subtraction. What fraction of the labile phosphate of the ATP has been converted into inorganic phosphate by the myosin? If desired, this experiment may be repeated at varying time intervals to establish the rate of enzyme action.
- 6. Preporotion of Muscle Fibrils (Perry). Rabbit muscle is dissected, chilled, and minced as for the preparation of myosin. The mince is homogenized in a Waring blendor for 2 minutes with 8 volumes of 0.08 M borate buffer, pH 7.1, at 0° C. The following operations are likewise carried out at this temperature. The homogenate is centrifuged 15 minutes at about 600 g., and the sediment is resuspended in the original volume of borate. This is homogenized once more for 2 minutes. This suspension is centrifuged for 20 minutes at 600 g.; the supernatant solution is discarded, while the lighter-colored upper layer of the sediment is taken up in fresh borate buffer. This suspension is then freed from coarser material by centrifugation for 5 minutes at 300 g. The suspension is then subjected to several cycles of centrifugation (20-30 minutes at 600 g.) and resuspension. Toward the end of the preparation 0.1 M KCI may be used instead of borate buffer.

Such suspensions form turbid viscous fluids which show a strong flow birefringeore. Microscopic observation reveals the presence of thin threads, in many of which the cross-striation is clearly visible. The addition of ATP results in rapid contraction, revealed by loss of birefringence and precipitation of floculent material. Fibril preparations are very active as ademosinetriphosphatase, but contain also myokinase and adenylate deaminase activity, the myokinase can be removed by washing ATP-contracted fibrils with water or dilute KCI solution

7 Contractile Fiber Preparation (Szent Gyorgyi) A rabbit is killed as for the preparation of myosin, and eviscerated The body is placed in a refrigera tor for 15 minutes and is finally cooled in ice. The psoas muscle is exposed by removing the sides and the anterior part of the body, and is liberated by placing the index finger under it and moving the finger in a longitudinal direction Then the muscle is lifted slightly and punched through with a probe or with small forceps with closed tips so as to separate a fiber bundle about 2 mm in diameter from the bulk of the muscle This fiber bundle is then separated over the whole length by moving the forceps parallel to the fibers A small stick (applicator rod) is placed alongside, and the fiber bundle Is tied to it at both ends, whereupon, it is cut free Other fiber bundles are Isolated in the same fashion Each bundle with its rod is placed in 50 per cent glycerol at 0°C (maintained with melting ice in a thermos bottle, placed in a refrigerator) The fluid is exchanged twice for fresh, precooled 50 per cent aly cerol, at dally intervals, after which the fibers can be stored at -20° C for months Before use, the fibers are placed in 15 per cent glycerol at 0° C for one hour, and are then split longitudinally into finer threads by pulling with fine forceps

Contraction can be observed by placing such fibers in fresh muscle extract (made by suspending freshly minced muscle in an equal volume of bolling water and filtering through cloth), or in 0 2 per cent neutralized ATP in 0 l Vi KCl, 0 001 Vi VigCl.

8 Preparation of Actin (Straub, with modifications of Bailey and Tiso) Nuscle is extracted for 10 minutes, and the mince is diluted threefold with water, as in the preparation of crystalline myosin. After straining through gauze, the residue is washed by gentle stirring in 4 volumes of 0.6 % I Nail Co. (the pil hecomes about 7) and then in 10 volumes of water, each washing lasta 20.30 minutes. The residue is pressed as dry as possible in a cloth, and is then extracted twice in a Waring blendor for I minute with 2 volumes of cold n butanol (let it stand for 30 minutes), and finally 3 times with 2 volumes of acctone, after which it is dried in the air.

The dry fiber is extracted with 25 volumes of water or (in order to assure a good jild) ATP solution (about 60 mg per liter, pli 8 2) The material does not wet casily, and must be thoroughly stirred for 30 minutes in the cold The solution is filtered by suction and centrifieded.

This extract contains actin in the globular form (usually about 5 mg per ml, ol about 50 per cent purity), so that its viscosity is low, and no biefringence is shown After the addition of 6 Lt to a final concentration of 0 1 M, rapid formation of fibrous actin takes place This is evident from an increase in viscosity, frequently leading to a gelation of the solution, and by the appearance of a most intense flow birefringence.

- 1. Extraction Procedure. About one g. muscle is extracted in the cold with 3 ml. of 0.5 N perchioric acid, and the residue (after filtration or centrifugation) is re-extracted with 2 ml. of the same reagent. The combined extracts are rapidly neutralized to a faint plac color with phenoiphthalein, and are left in the cold to assure precipitation of the potassium perchlorate (the residual solubility of which is about 0.05 M at 0° C.). The solution is filtered quantitatively into a 10 ml. volumetric flask, the phenoiphthalein is removed by extracting twice with ether (withdrawing the ether with an eyedropper), and the solution is brought to volume. Alternatively, neutralization can be controlled with the glass electrode.
- 2. Chromatographic Separations of Adenosinepolyphosphates (Cofin and Carter). A column is prepared (2 cm. height, 1 cm. diameter) of the anion-exchange resin Dower 1 in the chioride cycle³⁴ in a suitable chromatography tube. A muscle extract, prepared as described above, is diluted 5 times with water (in order to reduce the sait content, which would counteract retention by the column), the pH is adjusted to 9.5–10, and the solution is passed slowly through the column, which retains the nucleotides quantitatively. Elution is effected by successive application of 100–765-ml. portions of 0.03 N HCl; 0.07 N HCl; 0.02 M NaCl; and 0.01 N HCl, 0.2 M NaCl. These solvents clute AMP and IMP, ADP, and ATP, respectively. The cluates are collected in fractions of about 10 ml., in each of which the nucleotide is determined by measurements of the ultraviolet absorption, as described on p. 283 for ATP or, if no facilities for this are available, by the Bial test, p. 845.
- 3. Separation of Adenosinepolyphosphates by Paper Chromatogrophy. Smail amounts of muscle extract (e.g., 0.02 ml.) are applied to strips of suitable filter paper (e.g., Whatman No. 4); the size of each spot is kept small by intermittent application while drying after each delivery in a current of air. The chromatogram is developed by means of a solution of 60 volumes of n-propanol, 10 volumes of water, and 30 volumes of concentrated ammonia (Eggleton). Nucleotides can be located by observation in short-wave ultraviolet light by means of a Mineralight lamp. The experimental details of paper chromatography are described on p. 18.
- d. Determination of Free Creotine. For this and the following determinations, the tissue extraction should be carried out with especial care and rapidity. It is advisable to use 0.5 N perchloric acld which contains 20 per cent ethanol, cooled to -10° C., and to perform all manipulations at the lowest possible temperature until the extract is neutralized. In a 10-ml. volumetric flask, mir 1 ml. extract, a few ml. of water, 2 ml. of alkaline a-naphthol solution (0.5 g. freshig dissoived in 1 liter of stock alkali: 60 g. NaOH and 160 g. Na/CO, per liter), and 1 ml. of diacetyl (a freshig prepared 1:20 dilution of a 1 per cent stock solution). Compare the color or absorbancy (at 520 mm) with that given by a standard solution containing 0.005 mg. creatine treated in the same manner.
- 5. Determination of Free Inorganic Phosphote and of Phosphocreatine (Ennor and Stocken's adaptation of the method of Berenblum and Chain).
- (a) INORGANIC PROSPRATE. In a separatory funnel, place 2 ml. of neutralized muscle extract, 1 ml. of 0.25 N H₂SO₄ containing 4 per cent NaCl, and 1 ml.

²⁴ The resin is cleaned by alternate washings with acid and alkali, and is finally treated with 3 N HCl and washed with water until neutral.

of 0.5 per cent ammonium molybdate. Add these reagents in rapid sequence (5 seconds in total being permissible) and shake immediately for 10 seconds with 8 ml. of isobutanol. Alter separation of the phases (which is lacilitated by the salt added to the suilure acid solution), wash twice with 4 ml. of 1 N II.SOs, swirling but not shaking the contents of the funnel each time. In parallel experiments, run a hlank (2 ml., water) and a standard (2 ml., coatalning about 10 g. P.). The isobutanol phases are then shaken for one-half under the 1-12 ml. of stannous chloride solution (1 ml. of a stock solution of 10 g. SnCl; In 25 ml. of concentrated HCl, freshly diluted with 20 ml. 2N II.SOs). The aqueous layer is remnved, and the alcoholic layer is quantitatively transferred to a 10-ml. volumetric flask, the funnel is washed with ethanol, and the solution brought to volume with this solvent. Colorimetric readings are then performed at about 640 ml.

(b) INGRANG PROSPITATE + CREATERPHONDELTE. Place 1 ml. extract in a separatory funnel with 2 ml. 1 N H.SO, and 1 ml. 5 per cent molybdate, and keep this at room temperature for 15 minutes. Run a blank determination with 1 ml. water, and a standard with 1 ml., containing 10 µg. P. After it has stood, extract with isobutanol, wash with II,SO₄, and reduce with SnCl; as described for inoranic phosphates.

6. Demonstration of Phosphates and Magnesium in Muscle (Hürthle's Experiment). Tease a very small piece of frog's muscle on a microscope slide. Expose the slide to ammonia vapor for a few moments, then adjust a core flass, and examine the muscle fibers under the microscope. Note the large number of crystals of ammonium magnesium phosphate distributed everywhere throughout the muscle fiber, thus demonstrating the abundance of phosphates and magnesium in this type of muscle preparation (see Fig. 226).

7. "Fuchsin-frog" Esperiment. Inject a saturated aqueous solution of luchsin S into the lymph spaces of a frog two or three times daily for one of two days, in this way thoroughly saturating the tissues with the dye, Pith the animal (dinsert a heary wire or blunt needle through the occipito-atlantold membrane), remore the skin from both blind legs, and expose the scalar correl in one of them. Insert a small wire hook through the jaws of the frog and suspend the animal from an ordinary clamp or from ring. Pass electrodes under the exposed scalatio nerve, and after tying the other leg to prevent any muscular movement, atimulate the exposed nerve by means of make and break shocks from an induction coil. The stimulated leg responds by pronounced muscular contractions, whereas the tied leg remains inactive. Continue the stimulation until the muscles are fatigued. The muscular activity has caused the production of lactic actd and this in ture has recated with the injected luchsin to cause a plank or red color to develop. The muscles of the inactive teg still remain unchanged in color.

The color of the fuchsin S when fajected is red; this substance acts as a pilindicator, however, and is colorless at the pfi of normal tissues. Upon a timulating the muscles, as explained above, lactic acid is formed and the resultant lowering of pfi regenerates the original color of the dise.

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11

Nervous Tissue

In common with the other solid tissues of the body, nervous tissue contains a large amount of water. The percentage of water present depends upon the particular form of nervous tissue, but in all forms it is mariably greater in the gray matter than in the white matter. Embroone nervous tissues also contain a larger percentage of water than the tissues of the adult. The gray matter of the brain of the fictus, for instance, contains about 92 per cent of water, whereas the gray matter of the brain of the adult contains approximately 83 to 81 per cent. Adult whole brain (mixed gray and white matter) has an average water content of 77 to 78 per cent, human spinal cord contains about 75 per cent of water.

The solids of nervous tissue include proteins, lipides, extractives, and inorganic salts. In adult whole brain the relative amounts of these vanous components, in percentage of the total solids, are approximately as follows proteins 38 to 40 per cent, lipides 51 to 54 per cent, and extractive (including inorganic salts) 8 to 9 per cent. Other parts of the nervous 57

tem may show a somewhat different distribution

The Proteins. The proteins of nervous tissue which bave been failly well characterized include several globulins, nucleoprotein, and the but minoid neurokeratin Two of the globulins coagulate at 47° C and 70° to 75° C, respectively, the nucleoprotein coagulates at 56° to 60° C The relatively low coagulating temperature of one of the globulins of nervous tissue may be of importance in connection with the nervous manifesta

tions of heat prostration

The Lipides. Many types of hindes occur in nervous tissue Those which have been reasonably well characterized include (1) phospholipide or phosphatides, (2) cerebrosides or glycolipides, (3) sulfolipides or sulfatides, (4) gangliosides, and (5) cholesterol Other types will doubtles be recognized as progress continues in this field. It will be noted that the hipides are cliefly of the compound type, there is little or no true fat in nervous tissue. For example, of the 54 per cent of cerebrum solids which are composed of lipide material, the phosphatides constitute approximately 28 per cent cholesterol 10 per cent cerebrosides? For eent, and poorly characterized and similar material It has been shown that the relative amounts of certain of the lipides present in the brain may be in fluenced by age and sex

Phosi holdidle (Phosi hatides) The phospholipides or phosphatide include (1) 'ceithins, (2) cephalins, and (3) sphingomyelin.

Lecithins The lecithins are sometimes spoken of as "phosphorized fats" Their relationship to the fats is indicated by the formula of a typical lecithin

This lecithin is called oleyl-stearyl lecithin. On hydrolysis it yields oleic and, stearic acid, glycerol, phosphone acid, and choline Chemically, this compound may he regarded as an ester of choline and a phosphatidic acid, and is therefore a phosphatidyl choline. The phosphotic acids themselves contain fatty acids, glycerol, and phosphoric acid (hut no choline), and have been isolated from tissues as the calcium salts. There is some question whether phosphatidic acids exist as such in tissues, or are produced from lecithiis by enzymatic removal of choline during the treatment of the tissue.

There are different lecithms, depending upon the character of the fatty and radicals Most of the naturally occurring lecithms contain at least one unsaturated fatty and radical, but there are instances where both fatty and radicals are found saturated. The saturated fatty ands found include palmitic and sterric. The unsaturated ands are oleic, linoleic, linoleing and arachidomic (see Chapter 3). Representative types of the lecithm molecule have been synthesized in the laboratory, and in some instances found to be identical with material isolated from natural sources.

It is clear from the structure given above that two isomeric forms of a lecithin molecule are possible, depending upon whether the phosphoric acid-choline group is attached to the α (terminal) or β (middle) carbon of the glycerol residue, such lecithius are known as α - and β -lecithius respectively. There is no satisfactory evidence at present that β lecithins occur naturally

The lecthins are not confined to the nervous tissues but are found in nearly all animal and vegetable tissues, where they appear to be primary constituents of the cell Lecthin is soluble in chloroform, ether, alcohol, benzene, and carhon disulfide. It may be precipitated from chloroform or alcohol-ether solution by acctone Lecthin may be caused to crystallies in the form of small plates by cooling the alcoholic solution to a low temperature. It has the power of combining with acids, bases, and certain salts, such as cadmium chloride, and the hydrochloric acid salt forms a double salt with platinic chloride.

Pure levilim is colorless, it readily turns known on exposure to air, presumably because of oxidation of the unsaturated fatty ands present. The choline-phosphoric acid portion of the leathin molecule is highly watersoluble, the fatty acid portion is insoluble in water but soluble in fats Lecthin forms colloidal solutions in water which are of the hydroph her emulsoid type (see Chapter 1). If a portion of solid lecthin is placed in water and then observed under the microscope the lecthin will be seen diffuse out into the water in the form of long curving strands (myels forms) which bear a remarkable resemblance to the protoplasmic problemates of lower forms of life. The hydrophihe nature of the lecthin molecule may be of importance in connection with the structure as properties of cell membranes.

Choline is one of the products of hydrolysis of lecithin accounting to about 15 per cent of the molecule. It is trimethyl hydroxyethyl ammo num hydroxide, and has the following structure.

H₂C OH CH₂OH

Hac OH
Choline is a moderately strong base and fo

Choine is a moderately strong base and forms a crystallino double salt with platine chloride. In addition to its presence in the lecithin molecule, choine is found widely distributed elsewhere in nature either free of logical processes in combination with acetic and the compound acetyleholine is of significance in a reve activity as discussed later. In the detatached to introgen or sulfur) which is an important dietary requirement. Further aspects of the biochemistry of choline has been shown to be a source of the labile methyl group (methylment Further aspects of the biochemistry of choline will be found in

Cephalms The cephalms differ from the lecithins in both solubility and themical structure Differences in alcohol solubility are used to separate the lecithin fraction from the cephalm fraction is found to the state three types of compounds. Two of these types differ from the cithins in that the choline of the lecithin molecule is replaced by either the choline (colamne) CH₂OH CH₂NH₂ or serine CH₂OH CH M₂ ethanolamnic and phosphatdyl serine respectively. As with the lecithins the nature and type of fatty and residues joined to glycerol in the cephalms may vary and the problems.

alms may vary and the problems of isomerism are similar to those discussed above for lecithm

A third type of cephalm less well defined than the other two types acids mostiol as a characteristic hydrolytic product along with fatty sends phosphore acid ethanolamine and possibly galactose and tar taric acid. According to Poleit the mostiol containing phosphatide from the complex mostiol containing phosphatide from the more Woolley and named lipositiol. Further work is required to clarify this situation.

Lysoleothin and lysocephalm are prepared by treating lecithin or cephalin with cobra venom in enzyme present in the venom splits off

the unsaturated fatty acid radical leaving the structure otherwise unchanged Compounds of this type have a strong hemolytic action on red cells and the hemolytic action of venom is probably brought about in this way These lysophosphatides combine readily with cholesterol, molecule for molecule, the resulting compound having no hemolytic power

Sphingomyelin Sphingomyelin represents a third type of phospholipide found in nervous tissue. It is a diammomonophosphatide made up of a molecule each of fatty acid, phosphoric acid, and the bases choline and

sphingosine

Among the futty acids found have been the saturated acids—lignocene, $C_{24}H_{45}O_{2}$, and another acid probably hydroxysteane. The fatty acid is apparently in an amide linkage with the amino group of sphingosine. The complete structure would be as follows, R indicating the fatty acid radical

The content of sphingomyelin (and of leathin as well) is considerably increased in the liver and spleen in the condition known as Niemann-Pick's disease

ACETAL PHOSPHOLIFIDES (PLASMALOGENS) Phospholipides which un der suitable conditions give a positive test for aldehydes were first described by Feulgen and given the name plasmalogen These substances are found to contain fatty and aldehydes joined in acetal linkage to the glycerol of compounds having the basic leathin or cephalin structure

In this structure, R represents the remainder of a fatty acid aldehyde chain, and the introgenous base is ethanolamine. The crystalline acetal phospholipide isolated by Thannh user et al. I from beef brain contained.

¹ Than hauser Boncoddo and Schmidt J Biol Chem 188 417 (1951)

palmitic and steame aldehydes, glycerol, phosphone acid, and aminoethanol The presence of a serine-containing acetal phospholipide in brain has also been reported Nothing is known as yet concerning the

function or physiological significance of aectal phospholipides CERLBHOSIDES The cerebrosides (sometimes called glycolipides because of their carbohydrate content) are lipides containing carbohydrate one fatty acid, and sphingosine, but no phosphoric acid or glycerol Examples include phrenosine, kerasine, nervone, hydroxynervone Two general categorics are now known (1) galactohpides, containing galac tose as the component sugar, and (2) glucolipides, containing glucose Galactolipides are the more abundant form, glucolipides were originally identified in spleen from Gaucher's disease, a condition in which excessive amounts of ccrebrosides accumulate in the spleen, but are now known to be a normally occurring component of various tissues Cerebrosides appear to be an important constituent of the myelin sheath of medullated nerve fibers, which contain about three times as much cerebroside as is found in nonmedullated nerves. The fatty acid in phrenosine is the hydroxy acid phrenosime acid C2.H49(OH)COOH, and in kerasine it is lignocerie acid. The glycolipides are relatively less soluble in ether and more soluble in hot alcohol than the phosphatides

GANGLIOSIDES Gangliosides represent a class of cerebrosides first described by Klenk, which are found in brain gray matter, spinal cord and other tissues On bydrolysis they yield one molecule each of a base presumably sphingosine, stearie acid, neuraminic acid, and three molecules of galactose

SULFOLIPIDES (SULPATIDES) The sulfolipides are similar to the cerebrosides, but characteristically contain sulfurie acid esterified with cerebronic (bydroxylignoceric) acid Sulfolipides containing sphingosine, galactose, ecrebronic acid, sulfuric acid and potassium in equimolecular proportions have been characterized by Blix

CHOLLSTEROL Cholesterol, one of the primary cell constituents. 15 present in fairly large amount in nervous tissue. It is a monatomic alcohol containing one double bond with the empirical formula C27H45OH Its

structure is as follows

Cholesterol

The carbon atoms are numbered as shown for purposes of reference The four rings A, B, C, and D form the cyclopentano-perhydrophenanthrene nucleus, which is characteristic not only of cholesterol and other plant and animal sterols but also of a wide variety of naturally occurring compounds of surprisingly diverse physiological significance. These compounds include (a) the bile salts, (b) the steroid hormones, (c) the steroid vitamins, (d) the aglycone portion of the cardiae glycosides (digitalis, etc.), (e) the sapogenins, derived from plant saponins, and one might also include here the carcinogenic hydrocarhons of the phenanthrene type. The possible interrelationship of these varied compounds whose chemical structures are fundamentally similar, has attracted considerable attention, it has been shown for example that the cholesterol of the bile is a precursor of the chemically similar choic and derivatives also present

Cholesterol is soluble in ether, ebloroform, benzene, and hot alcohol it crystallizes in the form of thin colorless, transparent plates (see Fig 101) Cholesterol is present in ble, and occurs abundantly in one form of biliary calculus. It is also present in blood and its quantitative determination is of clinical importance (see p. 580). It has been found in feees, wool fat, egg yolk, and milk frequently in the form of its esters of higher fatty acids. It is found in the tissues of all animals. The cholesterol present in the animal hody has its origin in animal foods or synthesis in the hody, work with isotopes has shown that aceta each may he a major intermediate in this synthesis. It does not appear to arise from plant sterols. That cholesterol may be synthesized in the unimal hody is proved by experimental work using isotopes as well as hy such facts as that hens continue to lay eggs containing cholesterol when fed a cholesterol free duet and that the tissues of the rat contain the usual cholesterol content when fed a diet containing no sterols.

Various isomers and derivatives of cholesterol are found in plant and animal tissues only a fow may be considered here A further discus sion of steroid isomerism will be found in Chapter 26 Allocholesterol or coprostenol is an isomer of cholesterol differing only in that the double bond is in the 45 instead of the 56 position Coprosterol or coprostanol is formed by the addition of two hydrogen atoms at the double bond in allocholesterol It is found in the feces being formed by bacterial reduction Ergosterol (a provitamin D) differs from cholesterol in having two more double bonds, one in the 7-8 position and one in the side chain, with an additional methyl group in the side chain which thus becomes -CH(CH3) CH CH CH(CH3) CH(CH3) CH3 On sustable irradiation ergosterol finally yields calciferol with a rupture of the Bring 7-Dehydrocholesterol, found in the skin, is also activated by irradiation to produce a compound with vitamin-D activity Stigmasterol obtained from plant sources differs from ergosterol in having only one double bond in the nucleus (like cholesterol) and in having the second methyl group in the side chain replaced by an ethyl group The phytosterols or plant sterols comprise several different sterols According to Schoenheimer, the bulk of plant sterols do not appear to be absorbed by the body

Inorganic Salts. Nervous tissue yields about 1 per cent of ash which is made up in large part of sodium and potassium chlorides and phos-

phates The morganic salts of fresh brain and nerve tissue consist largely of sodium and potassium chloride, potassium being in excess over sodium. The content of sodium and potassium nons varies from one part of the brain to another, and has attracted considerable attention in relation to a possible role in incre action. The student is referred to Page's book for a further discussion of this subject (see Bibliography).

CHEMICAL CHANGES IN NERVOUS ACTIVITY

The carbohydrate metabolism of nervous tissue appears to follow much the same course as that of muscle tissue, going by way of pyruvate and lactate anacrobically, and the citric acid cycle aerobically. Since glycogen is low in amount in the brain and glucose appears to he the major car bohydrate substrate, this organ is more immediately dependent upoo the blood sugar supply. Nerves consume oxygen and liherate heat to a greater extent following stimulation.

The transmission of impulses along a nerve or from nerve fibers to muscle fibers or secretory cells or from oon erver fiber to another across synapses in ganglia, is thought to involve chemical changes, either directly or as the source of potential differences 'lecording to one view, the nerve action current results from a redistribution of diffusible ions between the center of the nerve and the periphery, taking place successively along the nerve fiber. This redistribution of ions may be under the cootrol of chemical acencies.

At the motor nerve endings to voluntary muscle, stimulation appears to liberate the compound acetylcholine (see below). Involuntary muscles and secretory cells are controlled by two sets of nerves sympathetic adparasympathetic. The parasympathetic fibers blorate acetyleboline at the nerve endings to such cells. Stimulation of such nerves causes smooth muscle fibers as in blood vessels, to relax with dilation of the vessels Stimulation of the sympathetic nerves brings about the liberation of at least two substances whose structures are known these are adrenaline (epinephrine suprarenine) and nor adrenaline(nor-epinephrine, arternol demethylated epinephrine).

Lsually only one of these two neurohormones as they are called, predominates in a particular organ or extract thereof. There is some basis for the view that nor adrenaline is a precursor of adrenaline. Of the two possible optical isomers of cach compound the leverotatory form is biologically much more active than is the dextrorotatory form

known as adrenerge nerves those releasing acctylcholine are called cholinerge nerves. The sympathetic fibers are adrenerge nerves, cholinergic nerves include motor nerves, all preganglionic fibers, and the postganglionic parasympathetic fibers.

Acetylcholine. As stated above, acetylcholine is believed to he the neurohormone of cholinergic nerves. Chemically, acetylcholine is the acetic acid ester of choline:

The structure as shown is that of the hase; the compound is usually obtained commercially as a halogen salt, the iodide heing less hygroscopie than the chloride or hromide. Solutions of acetylcholine salts show slight hydrolysis in neutral solution, which is accelerated in alkaline solution. Free acetylcholine may he detected, usually hy hioassay, in low concentration in nervous and other tissues. A relatively much greater proportion of the total detectable acetylcholine coutent of tissues is in a hound inactive form. The acetylcholine released hy nerve stimulation presumably comes from this source; the mechanism is not known.

Brain tissue contains an enzyme, choline acetylase, which promotes the synthesis of acetylcholine from choline and acetate, through the mediation of adenosinetriphosphate (ATP) and coenzyme A. This reaction is similar to other hiological acetylations involving coenzyme A (see Chapter 33), choline acting as acceptor of the acetyl group.

Physiological control over the excessive accumulation of acetylcholine constantly heing produced hy normal nerve impulses appears to he effected by the enzyme cholinesterase, which catalyzes the hydrolysis of acetylcholine into the relatively inert compounds choline and acetic acid.

$$\begin{array}{c} \text{CH}_1\text{CO O CH}_1\text{ CH}_2\text{N}\!\!=\!\!(\text{CH}_4)_1 + \text{H}_2\text{O} \rightarrow \text{CH}_3\text{OH CH}_1\text{N}\!\!=\!\!(\text{CH}_4)_1 + \text{CH}_4\text{COOH} \\ \text{Cholmesterase} & \text{OH} \\ \text{Acetylcholine} & \text{Choline} & \text{Choline} \\ \end{array}$$

Cholinesterase activity is found in the nervous tissues of all animal species thus far studied; it is also present in mammalian red blood cells, blood plasma, to a certain extent in other tissues, and in snake venom. The cholinesterase activity of the brain varies with location, the caudate nucleus heing much more active than other portions of the brain. The enzyme has not heen obtained in a pure or crystalline form; the most potent preparations have heen made hy Nachmansohn from the electric organ of the electric eel.

Study of cholinesterase activity in various tissues has revealed that cholinesterase is not a single enzyme but rather a name for a group of enzymes of varying characteristics and hiological significance. One classification groups the cholinesterases mto two broad classes, the "true" cholinesterases and the "pseudo" cholmesterases. The "true" enzymes are represented by the cholinesterases of nervous tissue, red

blood cells, and the electric organ of the electric cel and similar manne species. The cholinesterases of the blood plasma of most (but not all) animal species and of tissues other than nervous tissue belong to the class of "nseudo" cholinesterases

The two classes of cholmesterases are distinguishable from each other in terms of substrate specificity and hological occurrence and function. Thus, though both red cell ("true") and plasma ("pseudo") cholmesterase will act upon acetylcholine, other choline esters (propionylcholine, benzoylcholine) are much less readily hydrolyzed by the red-cell enzyme, which will, for example, rapidly decompose benzoylcholine. Distinction between the two types of cholmesterases is frequently based upon such differences in substrate specificity. Furthermore, the function of the cholmesterase of nervous tissue appears to be understandable in relation to the production and disappearance of acetylcholine in nerve function, whereas the plasma enzyme is one of the plasma proteins whose precise physiological function is not yet clear.

Interest in cholinesterases has been heightened by the discovery that drugs of the physostigmine (eserine) class, together with certain commercially important insectiedes, owe their mechanism of action to the ability to inactivate cholinesterase. Such anticholinesterase action can be demonstrated in vivo and in vitro. In the case of physostigmine, inactivation of the enzyme appears to be hased upon competitive inhibition, the drug competing with actylcholine for, and blocking it from, access to the enzyme molecule. This action is reversible in the laborator, by dialysis, enzymatio activity being restored when the drug is dialyzed away. In the case of patients, the cholinesterase activity of blood and tissies is lowered considerably after treatment with the drug, but returns to normal ia a matter of hours.

An anticholinesterase drug which irreversibly inactivates cholinesterase is illustrated by the substance dissopropylifuophosphate (DFP)

au enzymatically inactive product. Exactly how and why the DFP reacts with the polypeptide chain of the enzyme protein is not clear. Clinically, DFP is used in the treatment of glaucoma, paralytic ileus, and other conditions involving muscular dysfunction. Decrease in cholinesterase activity produced by DFP treatment in patients is much more prolonged than that produced by physostigmine, and essentially dependent upon the replacement of inactivated enzyme by new enzyme.

Insecticides whose mechanism of action is ascribed to anticholinesterase

activity include the following:

In the case of TEPP, insecticidal activity is due to a direct action on cholmesterase, TEPP being similar to DFP in this respect. Parathion does not have any significant anticholinesterase activity when quite pure; iusecticidal activity may be due to impurities, or to metabolic conversion of the P=S bond to P=O, which produces a substance (Para-oxon) with strong anticholinesterase activity. OMPA is another chample of a compound which has no anticholinesterase activity as such, but is converted by plant and animal tissues into a potent anti-cholinesterase, probably by oxidation within the cell, although the chemical nature of the reaction is not yet entirely clear.

EXPERIMENTS ON THE LIPIDES OF NERVOUS TISSUE

I. Preparation of Pure Lecithin. Free 4 pounds of brain tissue from adhering foreign tissue and mince in a chopping machine. Dry in a vacuum drying oven. (If such an oven is not available dry the tissue by treating several times with cold acetone.) Extract twice with acetone, using about 1.6 liters. Extract the residue with 2.4 liters of hot aicohol. Concentrate to one-third the original volume. Put in the refrigerator at 0° C. overnight. Filter. To the filtrate add a cold saturated solution of cadmium chloride sufficient to completely precipitate the lecithin.

To avoid decomposition in purification do not use high temperatures when effecting solution or concentration, and reduce the use of water to a minimum. The principal impurity in the cadmium precipitate is cephalin.

² The preparation from egg yolk is somewhat simpler. Stir the fresh egg yolks to a homogeneous emulsion with an egg-bester. Strain through cheesecloth. Pour into a double volume of hot 95 per cent alcohol. Allow to cool. Precipitate with cadmium chloride and purify as indicated above.

To remove this, shake the salts well with ether, and centrifuge. Repeat 8 to 10 times. Suspend 10 g of the salts in 40 ml. of chloroform, and shake at room temperature until a slightly opalescent solution is formed. Add a cold 25 per cent solution of ammonia gas in 95 per cent methyl alcohol as long as a precipitate is formed, avoiding a large excess. Centrifuge. Concentrate at about 10 to 15 mm pressure on a water bath at 35° to 40° C. Dissolve in drether and concentrate as before to get rid of moisture, Repeat. Dissolve in a minimum amount of ether and pour into acctone (about 50 ml.). Decant Dry the residue in a vacuum desiccator over H₁SO₄ and keep in a sealed glass tube to prevent oxidation.

2 Preparation of Crude Lecithin. Treat the finely divided brain of a sheep or a rabbit with enough ether to cover well, and allow it to stand in the cold for 48 to 72 hours. The cold ether will extract lecithin and cholesterol. Filter and add two volumes of acetone to the filtrate to precipitate the lecithin Filter off the lecithin, saving the filtrate for the preparation of cholesterol (see below) if desired. Test the lecithin as follows.

(a) Microscopical Examination Suspend a small portion in a drop of water on a silde and examine under the microscope. Do you see any "myelin forms'? (b) Osing Acto Ther 'Treat a small portion with osmic acid What happens? Why?

(c) ACEOLEIN TEST Make the acrolein test according to the directions on p. 106.

(d) Test for Phosphorus See p. 212.

3. Preparotion of Cephalin. Free 4 pounds of brain tissue from membranes, mince in a hashing machine, and dry in a vacuum drier. Pulverize and dry further. Extract exhaustively with acetone, using about 2 liters. Free from acetone in the vacuum drier and extract with about 3 liters of 95 per cent alcohol. Extract the residue with 2 liters of ether. Concentrate to a small volume Let stand at 0° C over night. Centrifuge and pour supernatant liquid into 98 5 per cent alcohol warmed to 60° C Dissolve the precipitate in ether Allow it to stand at 0° C over night. Repeat the precipitation with alcohol, redissolving in ether until on standing the ethereal extract no longer deposits a sediment of white matter. The final precipitate is dried and preserved as fol iecthin.

4. Preparation of Cholesterol* Place a small amount of finely divided brain tissue under ether and stir occasionally for one hour. Filter, evaporate the filtrate to dyness on a water bath, and test the cholesterol according to the directions given below. (If it is desired, the ether-acctone filtrate from simply necessary to evaporate she solution to dyness on a water bath.) Upon the cholesterol prepared by either of the above methods make the following tests.

(s) MICROSCOPICAL EXAMINATION Examine the crystais under the microscope and compare them with those in Fig. 191

(b) 1150, Test (Salaowski) Dissolve a few crystals of cholesterol in a little chloroform and add an equal volume of concentrated sulfuric acid. A play

⁴ The lecithin prepared in this way is satisfactory for the qualitative tests. Onne and serves to detect fata which contain unsaturated fatty and radicals—6.6 deep and in their indeed.

Furs cholesterol may be prepared from gallatones. Extract with benzene Evaporate

of colors from hluish-red to cherry-red and purple is noted in the chloroform, while the acid assumes a marked green fluorescence.

- (c) ACETIC ANHYDRIDE-H₅SO₄ Test (Lieaermany-Burchard) Dissolve a few crystals of cholesterol in 2 ml. of chloroform in a dry test tube. Now add 10 drops of acetic anhydride and 1 to 3 drops of concentrated sulfuric acid. The solution becomes red, then blue, and finally bluish-green in color.
- (d) FORMALDEHYDE-H₂SO₄ Test. To 2 ml. of a chloroform solution in a dry test tune add 2 ml. of formaldehyde-H.SO, solution (1 part of 40 per cent formaldehyde to 50 of the acid). Note the cherry color in the chloroform. Pour off the chloroform into another tube and add 2 to 3 drops of acetic anhydride. Note the hiue color. This test is said to he more delicate than Saikowskl's test.
 - 5. Preparation of Glycolipide. Mince 100 g. of sheep brains. Transfer to a 500-mi flask and add slowly with shaking 200 ml. of alcohol. Heat on a water hath for one hour with occasional shaking. Filter hot. Treat the residue a the water hath with another 100 ml. of alcohol for 15 minutes. Filter, and ombine the filtrates. Let stand over night. Filter off the precipitate on a mall paper. Transfer the paper to a heaker containing 50 ml, of hot alcohol. tir to dissolve the lipide and filter hot. Let cool. Filter or centrifuge, and vash the precipitate with ether to remove cholesterol. The residue consists namly of the glycolipides phrenosin and kerasin.
 - (a) Microscopical Examination Suspend a small portion in a drop of water
 - on a slide and examine under the microscope. (b) Solubility Try the solubility of the glycolipide in water, and in dilute
 - acld and alkail, and in hot and cold alcohol and hot and cold ether. (c) Phosphorus Test for phosphorus according to the directions on p. 212.
 - How does the result compare with that obtained with lecithin? (d) Place a little giycolipide on platinum foil and warm. Note the odor.
 - (e) Hydrolysts or Glycolymbe Place the remaining glycolipide in a small evaporating dish, add equal volumes of water and dilute hydrochloric acid, and boil for one hour. Cool, neutralize with solld sodium carhonate, filter, and test with Benedict's solution. fs there any reduction, and if so how do you explain it?
 - (a) ROSENIEDI'S PERIODIDE TEST Prepare an alcoholic extract of the fluid 6. Tests for Choline. under examination, and after evaporation apply Rosenbeim's iodo-potassium iodide solutions to a little of the residue. In a short time, dark brown plates and prisms of choline periodide hegin to form and may be detected hy means of the microscope. Occasionally they are large enough to be visible to the naked eye. They somewhat resemble crystals of hemin (see p. 483). If the slide he permitted to stand, thus allowing the fluid to evaporate, the crystals will disappear and leave brown oily drops. They will reappear, however, upon the addition of fresh lodine solution. v. Stanek claims that this
 - choline compound has the formula C.H., NOI Is. (b) ROSENIEIM'S BISMUTH TEST Extract the fiuld under examination with absolute alcohol, evaporate, and re-extract the residue. Repeat the extraction several times. Dissolve the final residue in 2 to 3 ml of water and add a drop of Kraut's reagent. Choline is indicated by the appearance of a bright brickred precipitate.

⁴⁻⁷ See Appendix.

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12

Enzymes and Their Action: Cell Respiration

The myriad chemical transformations going on continually in living matter would not be possible without enzymes, which are the most important tools of the living cell For the hydrolysis or the oxidation of such substances as fats and proteins, in the laboratory we commonly employ strong acids or alkalies or oxidizing agents and high temperatures Such reagents and temperatures are incompatible with the existence of living matter The cells are, however, able to carry out such reactions at approximate neutrality, at body temperatures, and at high speed with the aid of enzymes As an example it has been shown that it takes 10,000,000 times as great a concentration of hydrogen ion as of the enzyme sucrase to decompose a given amount of cano sugar in a given time at hody temperature

Enzymes are organic catalysts produced by living organisms. They are generally soluble and colloidal substances, characterized by great activity, specificity and susceptibility to the influence of pH, of temperature, and

of other environmental changes

Catalysis As indicated, enzymes act catalytically Catalysts are substances that alter the speed of chemical reactions without themselves undergoing any permanent change This definition indicates that cata lysts do not initiate chemical reactions but only speed up reactions already proceeding at a slow rate Thus hydrogen perovide undergoes a very slow spontaneous decomposition at room temperature with the formation of water and oxygen The addition of a little finely divided platinum or of an enzyme called eatalase enormously mereases the rate of this decomposition Catalysts function in many reactions, however, which cannot definitely be shown to be proceeding in their absence. It may be assumed in these cases that the reaction is actually proceeding but at an immeasurably slow rate From the practical standpoint, however, the catalyst or enzyme in these cases does initiate a reaction

Inorganic catalysts are of great importance in industry Thus sulfur diovide reacts slowly with ovygen to form sulfur triovide If, however, oxides of introgen are present, they catalyze the reaction so that it proceeds much more rapidly, eg,

$$NO + \frac{1}{2}O_{2} = NO_{2}$$

 $SO_{2} + NO_{3} = SO_{3} + NO$

Further in alogies between enzymes and morganic catalysts will be men tioned later 303

Classification. All the enzymes which have been isolated in pure condition to date are proteins. The properties which are used to classify proteins are much too general to be of use in classifying enzymes. Hence it is customary to classify them according to their action. Some enzymes contain characteristic prosthetic groups conjugated with the protein, and these groups are also used for further classification. For example, exidizing enzymes are commonly subdivided according to whether they contain hence, riboflavin, etc., as prosthetic groups. A considerable number of enzymes can be classified under the general term transferases since their action is to transfer such groups as amino, methyl, accetyl, glucosyl, fructosyl, or phosphoryl from one compound to another.

Hydrolasis. A great many enzymes catalyze hydrolysis reaction. This large group is commouly designated as hydrolases. The hydrolases, like the countries of their substrates, i.e., the substances on which they act. The nomenclature is harmonized as far as possible by using the termination ase in the names of enzymes. Thus we have esterases hydrolyzing estera, carbohydrases hydrolyzing carbohydrates, proteinases hydrolyzing proteins, amidases hydrolyzing amides, ctc. Under each of these groups are individual enzymes which act on specific substrates, such as maltase which splits maltose, lactase which hydrolyzes lactose, etc. The source of the enzyme may also be indicated in its name, such as salivary amylase, pancreatic lipase, etc. Some of the old names for enzymes are still used as a matter of convenience—such names, for example, as pepsin and trypsin.

ONDEANG ENDAMS Another large group of enzymes catalyzes oxidations. These may be called oxidizing enzymes. Oxidizing enzymes are frequently divided into two groups—oxidases and deby drogenases—which are distinguished by their mode of action. This is not an entirely satisfactory divisions since some enzymes can be placed in either group. A monsuitable classification makes no attempt to distinguish between oxidase and debydrogenases—other enteria are used instead. For example, the following groups may be recognized: (1) Enzymes containing into, (2) enzymes containing copper, (3) enzymes containing eoenzymes I of II, (1) enzymes which entally are the reduction of cytochrome, (5) enzymes which contain riboflavin () ellow enzymes), and (6) various other oxidizing

There are still other enzy mes that do not fit into either of these groups for example, phosphory lases act similarly to hydrolases, except that a phosphoric acid residue and not water, is used in splitting the substrate. There are other enzy mes the hydrases, which remove or add water to the substrate without hydroly zing it, and there are still other enzymes, the time desimolases which catalyze the splitting of a carbon chain. Thus, any mohixase catalyzes the reversible splitting of fructose-1,6-diphosphate into the throsphosphates dhydroxyactone phosphate and phosphoglycare aldchyde. The mutax's catalyze the oxidation of one molecule of the substrate and the simultaneous reduction of another molecule of the substrate. Recently aldchyde mutax has been shown by Rucker to be 1 othing more than a mixture of alcohol dehydrogenase and aldchyde dehydrogenase.

In a number of cases, two or more names are commonly applied to the same enzyme For example, amyluse is ulso known as diastase, sucrase is commonly called invertase or saccharase, cytochrome oxidase was formerly called indophenol oxidase, etc Sometimes such synonyms arose because enzymes from different sources at first were not recognized to be identical For example, tyrosuase or monophenoloxidase was at one time thought to be distinct from polyphenoloxidase. In other cases the original name given to an enzyme has proved to be inadequate For example, the name zymase was at one time used for the yeast enzyme which ferments sugars to alcohol. Subsequently it was found that the original zymase was really several enzymes and the original name had to be discarded.

A list of some of the more important enzymes is given in this connection, without any attempt to adhere to a rigid classification (see Table,

pp 306-307)

Chemical Nature. The problem of the purification of enzymes has been essentially one of their separation from other associated proteins Because of the difficulties of such separation and purification, the chemical nature of enzymes has been much in doubt. The first preparation of a erystalline enzyme was that of urease hy Sumner in 1926 (see Fig. 75) Since that time crystalline pepsin was obtained by Northrop (see Fig. 76), crystalline trypsin and chamotropsin by Northrop and Kunitz, and pepsinogen by Herriott and Northrop Crystals of yellow respiratory ferment were obtained by Warhurg and those of β -amylase (Fig. 74) by Balls To date, about 60 enzymes have been obtained in crystalline form However, a few of the claims made are either erroneous or duhious All of these crystalline preparations have proved to he proteins, and the specific action of these enzymes has never been observed in the absence of the specific proteins. There is additional evidence which shows rather clearly that the enzymes and erystalline proteins are identical If the protein is destroyed or denatured, the enzyme action is lost Also, various crystalline enzyme preparations have been tested for homogeneity, using such criteria as diffusion, movement in an ultracentrifuge migration in an electrical field, solubility, etc , and with the exception of pepsin each has been found to consist of a single component Pepsin appears to be a mixture of similar proteins. Some of these crystalline enzymes have heen recrystallized as many as ten times without change in composition and properties

Further evidence that the enzyme itself is a protein has been presented, as in the case of pepsin, by showing that if the protein of the material is denatured through the action of alkah, the activity of the enzyme is decreased in a parallel manner. For example, pepsin becomes completely denatured and mactive at pH 105. If the solution is adjusted to pH 5 is and is allowed to stand at this pH, a small part of the protein is renatured and there is a corresponding appearance of activity of the enzyme. The solution should be adjusted to about pH 1 before testing for the enzyme activity.

Crystalline trypsin denatured by heat also shows on standing a parallel reappearance of activity and of native protein in the solution. The trypsin

CLASSIFICATION OF ENZYMES

hame and Class	Dud shulson	Substrate	End-products	
Hydrolases Ca bohyd ases 1 Amylase	Pancreas anava malt.	Carboby drates Starch, destr n etc	Maltose and destrins	
2 Lactase	intest nal pure and	Lactore	Glucose and galactese	
3 Maltase	muccosa Intest nal purce 3 cast	Maltone	Glucose	
4. Sucrase	Intest nal juice, yeast	Cucrose	Glucose and fructose	
o Emulsin	Plants	B-Glucoudes	Glucose etc	
\ucleases		Nuclese acid and de-		
1 Polynucleot dase	Panereat cauce nuc-	Nucle c acid	\ucleotides	
2 Nucleot dase	tinal ju ce, etc Intest nal juce, and other tuoues	Nucleotides	Sucleondes and plos phone and	
3 Nucleosidase	An mal tusues	Nucleonides	Carbohydrate an	
Amidases 1 Argmase 2 Urease	Liver Bacteria, soj bean, jack bean, etc	Amino compounds and an idea Argin ne Urea	Orn thine and ures Carbon dioxide at animonia	
3 Glutaminase	Liver etc	Glutamune	Glutamie acid and ar	
4. Transaminase	Animal tunues	Glutame and and exal- acet e acid ete	monia a-helogistane and a part e acid, etc	
Purine Deaminases		Pumpe bases and deny		
1 Adenase	Atumal tunues	Adenine	Hypoxanthine and at	
2 Guanase	An mal t sauce	Guanine	Nanth ne and ammor	
Peptidases 1 Aminopoly peptidase.	least, numi nes etc	Peptudes Pulypept des	myler peptides at	
2 Carboxypepudase	Panereas	Polypeptides	amino acida	
3. D peptidase	Plant and animal to		Amino ac da	
4 Prokusse	Animal tustiers and Jesst		Proline and simp	
Pro cinases I Pepein		Proteins		
	Gastrie juice	Prote as	Proteoses pepton	
2. Trypsin	Pancreatic juice	Protesus, proteoses and	Polypept desand am	
3 Cathepan 4. Rennin	Atomal tosues	Proteins	Proteoses and peptor	
5 Chymotrypan	f all stomach			
6 Papain	Pancreat e itisce	Proteins, proteores, and	Polypertides and am	
7 Fein	Papaya, other plants	Proteins, protecess and	Polypertides and ami acids	

CLASSIFICATION OF ENZYMES-(Continued)

CLASS			
Name and Class	Distribution	Substrate	End products
Malic deby drogenase I Lactu deby drogenase J Lactu deby drogenase J Lactu deby drogenase J Lactu deby drogenase J Lactu deby drogenase J Lactu deby drogenase J Lactu deby drogenase J Lactu deby drogenase J Lactu deby drogenase J Lactu deby drogenase J Lactu deby drogenase J Lactu deby drogenase J Lactu deby drogenase J Lactu deby drogenase	namal and I lant to auces minal and plant to- ninal and plant to- ninal tessues and jeast hidneys and heart there's and heart through the subsection of the	Ett 31 alcohol and other alcohols (2-) Male seed Lectic acid Lectic acid Lectic acid c. 3 Hydroxybut; ric acid occlicose Robison extra thexoe-d-phosphate) (b) ecrophosphate Aldebydes	Oxalosuccinic acid Oxalosuccinic acid Pyruvic acid
Rnzymez II hich Reduce Cy.oci rome 1 Succinic deh) drogenase (as ordinaril) prepared)	Plants animals and	1 Successive need	Fumaric acid
1 ellow Ensymes 1 Warburg a old yellow ensyme 2 Diaphorase 3 Hass ensyme 4 Nanthine oxidase 5 D-Amino acid oxidase 6 L-Amino acid oxidase 7 TPN Cytochrome Creductase 8 DPN Cytochrome Creductase 1 Funarase 1 Funarase	Living organ sms	Reduced coensyme II Hypoxanthine xer thine aldehydes re duced coensyme ete p-Amino acids + Or	ensyme ure acid antime material entered even an acid and acid even an acid and acid entered even and acid acid and am constant entered even acid acid and am constant entered even acid acid acid am constant entered even acid acid acid am constant entered even acid acid acid am acid
2 Acomtase 3 Enclase		and 2 Phosphogly certe a	t-isocitric ac d Phosphopy ruvio ac d + H ₂ O
Mutases 1 Glyoxalase	Laung organisms general	in Methyl glyoxal a other aubstituted a oxals	rly
Desmolasea 1 Z) moherase (aldola: 2 Carboxylase 3 \$\beta\$-keto carboxylases 4 Amino neid decarb ylases 5 Carbonic anhydras	Plant tissues Animals bacte plants Plants animals	bac- L Amino acids Carbonic ac d	phosphoglycera act Acetaldebyde and CO c-heto seids Amines and CO: GO: + H:O
Other enzyroes 1 Phosphoralise 2 Phosphohexouson e 3 Hexokinase 4. Phosphoglucomut.	Yeast, annual to	t t = Glucose 6-phosphs Adenosinetriphosp	hate Fructuse 6-phosphate Adenosinediphosphate + glucose 6 phosphate + phate
may also be hydrolyzed hy pepsin in acid solution, and the loss in activity

is just proportional to the hydrolysis of the trypsin protein.

In the case of those crystalline enzymes which have been most studied it seems clear that the enzyme activity is bound up with the integrity of the protein of the preparation and that these enzymes are thus properly

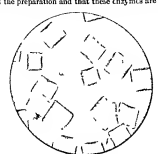


FIG 74 β-AMYLASE CRISTALS Courtesy, Dr A. L. Balla.



FIG 75 UREASE CRYSTALS. Coursesy Dr J B Summer

FIG 76 PEPSIN CHYSTALS. Courtery Dr J IL Northrop.

characterized as proteins. Experiments by Northrop on the acetylation of pepsin suggest that the activity is bound up not with the primary amino groups, but with the presence of free hydroxyl groups of tyrosine. That certain cusymes may one their activity to groups of a nonprotein character combined with protein is clear from our knowledge of the outdasses, some of which appear to be hime-protein derivatives, the activity

of which is bound up with the iron-pyrrole group Perovidase is a specific example of such an oxidase Heme has, however, much less peroxidase activity than its natural protein combinatious, so that both elements are evidently essential for proper function Preparations of certain other enzymes—such as amylase, sucrase, and lipase—have been variously reported as protein-free or very low in protein Protein tests made on enzymc solutions are, however, inconclusive, since the protein tests are far less sensitive than the tests for enzyme activity As to the nature of the groups in the enzyme molecules which are responsible for the specifiety of their action, almost nothing is known except in the cases of some of the oudases

Specificity and Mechanism of Enzyme Actions. Not only do protemases not act upon carbohydrates, nor earbohydrases upon proteius, hut much greater degrees of specificity exist Thus maltose, lactose, and sucrose require different enzymes for their hydrolysis Further, the maltase of yeast splits maltose (an α -glucoside) and certain other α glucosides, but not β -glucosides, but β -glucosidase or emulsin hydrolyzes only β -glucosides The maltase will not hydrolyze any glueosides in which the a-glucose residue has heen altered The specificity appears to depend primarily on the glucose residue and secondarily on the group attached to it The dependence of enzyme action on the configuration of the substrate is further brought out by the fact that dipeptidase will attack glycyl-1-leueine but not glycyl-D leueine It was findings of this sort that led Emil Fischer to suggest that the relation of euzyme to substrate was much like that of a key to a lock

It has been pointed out that enzymes are eatalysts Our knowledge of the mechanisms involved in enzymic catalyses is, however, very slight There is much evidence that enzymes form intermediate compounds with the substrates on which they act This intermediate compound is believed to be much less stable than the original substrate so that the latter now breaks down spontaneously, the enzyme being again liberated According to the views of some workers there may be two groups (or perhaps more) in the enzymo which are involved, one group acting chiefly to bind the enzyme with the substrate while the second combines with another part of the substrate so as to reduce the stability of the substrate This view has been developed especially in connection with the peptidases

Intestinal dipeptidase will split all peptides made up of naturally occurring amino acids. It will not spht peptides containing the optical isomers of these amino acids Thus glycyl-Leucine will be split but not glycyl-Deleucine If the free earboxyl group of the dipeptide is esterified (as by making the ethyl ester), hydrolysis still occurs, indicating that the enzyme does not combine with the carboxyl group If the free amino group is acetylated, the compound is not split This indicates that the enzyme combines with the amino group of the peptide If the H of the NH group of the peptide is substituted, it is not acted upou Replacement of the H

of the adjacent C atoms also prevents splitting These observations indicate that the enzyme must have at least four groups showing a similar space relationship to these four groups in the dipeptide It has been suggested further that the enzymo combines with the dipeptide first through the amino group and then a second group in the enzyme attaches to the NH group, this latter linkage leading to a lowered stability of the peptide linkage so that splitting occurs. It has been shown that acetylation of the NH group in peptides does lead to instability. The findings may be summarized diagrammatically as follows it must be understood, however, that no explanation of the mechanism of such enzyme actions is yet established.

Kulin and Hartree observed that peroxidase forms compounds with hydrogen peroxide that show definite absorption bands. Chance has investigated these compounds also compounds of catalase with hydrogen peroxide. The reaction of catalase with hydrogen peroxide is

(1)
$$Cat(OH)_4 + H_2O_2 = Cat(OH)_4OOH + H_2O$$

(2) $Cat(OH)_4OOH + H_2O_2 = Cat(OH)_4 + H_2O + O_2$

The reactions with peroxidase would be
(1) PerOII + II₂O₂ = PerOOH + II₂O

(2) PerOOH + AH₂ (oxidizable substance) = PerOH + H₂O + A

Action of Physical and Chemical Agents on Enzymes. Variationof temperature and of hydrogen ion concentration of the medium have
a marked effect on enzyme action. The speed of action must naturally also
vary with the concentration of the enzyme and the concentration of the
substrate. The effects of activators, of coenzymes, and of a proper coacentration of electrolyte are of importance in many cases, as well as the
intuitive effects of inhibitors or "poisons". As digestion proceeds, the
products of the reaction may have an inhibiting effect

Influence of Temperature Most enzymes are mactivated by beating their solutions to temperatures in the neighborhood of 60° C. This mactivation is associated with a denaturation of the protein of the preparation. In some cases there is, on standing an appreciable reversal of the denaturation process with the reappearance of some enzyme activity. Heat resistance may be influenced by the presence of protective substances, such as the enzyme substrate, or by pH. Trypsin is more resistant in acid than in alkalinic solutions and is in fact very resistant. In acid solution it may endure temperatures near the boiling point. Under these conditions it is denatured but the process is rapidly reversed on cooling

reinperatures as low as freezing do not commonly destroy enzymes. Within their active range most enzymes have a temperature coefficient (Qio) of about 2 i.e. they approximately double their activity for each 10 degrees (Tise in temperature As the temperature rises, however inactivation influences the result—since inactivation also mercases with

rise in temperature. The optimum temperature for the action of an enzyme is thus determined by the balance between nucreuse of activity with temperature and decrease in amount of enzyme present due to mactivation In prolonged experiments enzyme destruction becomes more important than in short experiments, and the optimum temperature for such experiments is therefore lower The majority of enzymes act most rapidly at 40° to 50° C. For certain plant protein ases the optimum is

Influence of Hydrogen-ion Concentration. Enzymes are mactihigber vated by certain degrees of acidity or alkalinity Thus yeast sucrase is inactivated rapidly below pH3 and pepsin above pH8 By the optimum pH of an enzyme is meant the pH at which it shows greatest activity For most enzymes this point has between pH 4 and 8, and for many between pH 5 and 7 For practically all enzymes the pH range is rather narrow and activity falls off rapidly on both sides of the optimum The best pH depends to some extent on temperature and enzyme and substrate concentrations The nature of the buffer may ebange the optimum pH of an enzyme considerably The same is true for the nature of the substrate When the substrates are ionizable, as in the case of proteins the optimum pH may vary with the substrate This is true for pepsin. One of the lowest pH optima is that for pepsin (pH 2) and one of the bigbest that for blood phosphatase (pH 9) Alterations in pH may affect the dispersion of the enzyme, the rapidity of combination of enzyme and substrate, and the decomposition of the enzyme-substrate compely with the formation of reaction products

Influence of Other Physical Agents Enzymes may be destroyed by sbaling by ultraviolet irradiation and by v rays These agents probably

act by denaturing the enzyme

Influence of Chemical Agents Most enzymes when acting at optimum temperatures will show, on the addition of chemical agents either an inhibiting effect or no effect at all In some cases, however, an acceleration is noted, and certain enzyme preparations are mactive unless additions are made

If salivary amylase is dialyzed free from NaCl it becomes mactive on starch but activity is restored on the addition of sodium, potassium, or certain other chlorides. The chloride here may be called a cocnzyme 1e, it belongs to a class of substances which are specific for particular enzymes and necessary for the activity of those enzymes. In this sense phosphate which is essential for the action of zymase, is also a coenzyme as is the organic crystalloidal substance cozymase which is also necessary for zymase action More recent usage limits the term coenzyme to organic compounds which act in this manner, e.g., diphosphopyridine nucleotide.

Many enzymes are ictivated by such metallic ions as Mg++, Co++,

Mn++ Zn++, etc Emil Smith bas studied the activation of various peptideses and believes that at least two of the honds between substrate and enzyme are through the metallic ion

Activators and kimases are believed to hring about chemical changes in the inactive forms of enzymes, so as to render them active. Thus HCI changes inactive pepsinogen to active pepsin. The pepsinogen here is called a zymogen or proenzyme and the HCI (or more strictly the H ion) is spoken of as an activator. Activators of an organic character are sometimes called kinases. Thus trypsinogen is changed to active trypsin by enterokinase. Enterokinase is prohably an enzyme. It can be replaced by trypsin the action of trypsin upon trypsinogen, therefore, is autocatalytic Chymotrypsinogen and procarboxypeptidase also are activated by trypsin.

Enzymes are possoned by a variety of substances which form compounds with them A number of heavy metals stop enzyme action, apparently by combining with the enzyme acting as an acid, though combination with sulfhydryl groups in enzymes may also he involved. Urease is sensitive to traces of heavy metals. It is may combat this toxicity of

the heavy metals probably by combining with them

HCN and H.S have little effect on most hydrolytic enzymes. The protenase called papain is, however, activated by them, and the action of kcratinase is much facilitated by the presence of H.S. Certain oxidases are mactivated by these substances in small amounts, apparently because they combine with the active iron of these enzymes.

Certain acids, such as phosphotungstic or pieric acid, combine with the

enzyme, acting as a base, to form mactive compounds

Antienzymes include various naturally occurring inhibitors with some degree of specificity. The blood serum contains antitrypsin, and the mucosa of the intestinal tract appears to contain some antipepsin and antitry pair. In addition there are various antienzymes which are true immune bodies produced by the usual methods for producing antibodics. For example, antiluciferase, antiurease, and anticatalase have been produced by in jecting the appropriate enzymes into animals.

Linetics of Enzyme Action. Enzyme kinetics deals with the velocity of enzyme reactions and with agents or factors which affect this velocity

It is usually possible to cause the velocity of an enzyme-catalyzed reaction to remain constant for a sufficient period of time to allow measurement of the enzyme by determining the reaction rate, if enough substrate is used and if other factors with an II.

urement of the enzyme by determining the reaction rate, if enough substrate is used and if other factors such as pH, temperature etc are held reasonably constant 1 reaction of this variety is called one of apparent zero order and follows an equation of the form

T

(1)
$$\frac{x}{t} = I \quad \text{or} \quad x = It$$

where x is the amount of substrate decomposed in time t, $\frac{x}{t}$ is the reaction velocity, and t is a constant depending upon the amount of enzyme present. Provided that factors such as pH and temperature are held constant,

I can be used as a measure of the amount of enzyme present

If the amount of substrate employed is low, as is necessary with enzymes such as catalase which are gradually destroyed by excess sub-

strate, the reaction velocity is often proportional to the amount of substrate present at any time Such a reaction is called an apparent monomolecular or first-order reaction and follows the equation

$$v = \frac{dx}{dt} = k(A - x)$$

(deduced from simple application of the mass law) where the velocity v must be expressed using the calculus notation $\frac{dx}{dt}$ instead of $\frac{x}{t'}$, since v varies continuously with x. The term A signifies the initial concentration of substrate, x is the decrease in concentration of substrate (equivalent to the increase in concentration of products) that has occurred at time t, and t is a constant which varies with the amount of enzyme (at constant pH, temperature, etc.) and can be used as a measure of the amount of enzyme present

Since Equation (2) is not applicable as it stands, it must be transformed by use of the integral calculus to eliminate the $\frac{dx}{dt}$ term. When this is done the following equation is obtained

(3)
$$\log \frac{A}{(A-x)} = kt$$
, or (3a) $k = \frac{1}{t} \log \frac{A}{(A-x)}$

If it is desired to obtain the theoretical initial reaction velocity, this can be done by multiplying L by A, or

$$v_* = kA$$

as is apparent from an inspection of Equation (2)

The use of the term apparent in describing the order of enzyme-catalyzed reactions is necessary because such reactions usually consist of a sequence of himolecular reactions, one of which may be rate-limiting and therefore control the observed kinetics. In case water is involved in a rate-limiting himolecular step, a truly himolecular reaction will follow monomolecular kinetics, as is explained in texts on physical chemistry, owing to the essentially constant water concentration. Bimolecular kinetics are observed with only a few enzymic reactions and will not be considered here.

In case the concentration of substrate is neither very high nor very low, the observed kinetics of an enzyme-catalyzed reaction may not follow either an appyrent zero-order or an apparent first order reaction. In such a case it is still possible to obtain a valid measure of the amount of enzyme present from kinetic studies if one can measure the length of time necessary to att iii decomposition of a certain fixed percentage of the substrate (say 5 per cent). In this case the amount of enzyme is proportional to the respocal of the time value.

As has been inferred above, the rate of an enzyme-catalyzed reaction

can be affected by factors such as change in temperature, change in pH, and change in substrate concentration (or partial pressure, if a gas is reacting) In addition the enzyme may be mactivated by the substrate or

inhibited by various types of enzyme inhibitors

Temperature change generally results in a bell-shaped curve when enzyme activity is plotted against temperature, with a pronounced optimum The latter results from an enhancement of reaction rate as the temperature increases, followed eventually by a decrease in rate owing to the denaturation or destruction of the enzyme at higher temperature values The exact position of the temperature optimum is not of theoretical significance however, since it is dependent upon various factors such as substrate concentration, pH, and the time interval during which the measurement of activity is carried out. It is very difficult to arrive at a good estimate of the initial reaction velocity in determining temperature optima Hence a somewhat better procedure is to determine the effect of temperature on the enzyme by treating it at various temperatures for a given length of time and then determining activity under standard conditions of pH, temperature, and substrate concentrations By this procedure an S-shaped curve is obtained which gives a more reproducible measure of the destructive effect of temperature on the enzyme The effect of temperature on enzymatic activity is of theoretical significance (in calculating energy of activation) in the range where no damage to the enzyme is occurring, and throughout this range enzyme activity generally increases with temperature increase in a predictable way

Curves showing the effect of pH on enzyme activity in general give optima which are of theoretical significance. Changes in pH may affect primarily the substrate or the enzyme, or both, and are concerned with producing the necessary ionic states of the enzyme or substrate or both

A fundamental approach to enzyme kinetics was developed by Yichaelis and Menten?, who proposed to measure the affinity of an enzyme for its substrate by studying the effect of substrate concentration on initial reaction velocity with a constant amount of enzyme present. The theoretical treatments is based upon the assumption of the following sequential reactions involving enzyme, substrate and reaction products.

(4)
$$E + S \stackrel{k_1}{\underset{k_2}{\rightleftharpoons}} ES \stackrel{k_2}{\sqsubseteq} E + P$$

where E stands for enzyme S for substrate and P for products. The L's are the rate constants for the reactions as indicated by the arrows

In the simplest case it is assumed that the rate-limiting reaction is $ES \rightarrow E+P$ the formation of ES being much more rapid. By a simple algebraic process if this assumption is made, the following equation can be derived, which shows the relation between the maximum reaction velocity $V_{\rm es}$, obtained when an excess of substrate is present and the enzyme is saturated with substrate and the reaction velocity $v_{\rm e}$ obtained with substrate concentrations below the saturation level

^{*} Michaelis and Menten Biochem Z 49 333 (1913)

(5)
$$v_{i} = \frac{V_{m}[S]}{[S] + K_{m}} \text{ or solving for } K_{m},$$

(5a)
$$K_m = [S] \frac{V_m - v_*}{v_*}$$

where v, is initial reaction velocity for the enzyme-catalyzed reaction in question, [S] is substrate concentration, and Km, the Michaelis constant, is the dissociation constant for the enzyme substrate complex 1e, the equilibrium constant for the reaction $ES \rightleftharpoons C + S$ In this simple case therefore, $K_m = \frac{k_2}{l}$ where the small l's are defined as above. In ease the decomposition of the ES complex to yield enzyme + products is not strictly rate-limiting, K_m must be interpreted as $\frac{k_2 + k_3}{L}$. It will be noted that in the simplest case Km is an inverse measure of the affinity of enzyme for substrate 1e the smaller the Km value, the higher the affinity

In theory, K_m is equal to the substrate concentration when $v_i = \frac{1}{2}V_m$ However, in determining K, values Equations (5) and (51) are not very suitable, since Vm is not easily obtained Equation (5) is a branch of an hyperbola, but in practice if an attempt is made to obtain Vm from an estimate of the horizontal asymptote, large errors usually occur for various reasons However, it is possible, as shown by Lineweaver and Burk, to transform Equation (5) into a linear equation simply by equating the reciprocal of both sides and simplifying by division Equation (6) is thus obtained

(6)
$$\frac{1}{v_s} = \frac{1}{V_{ee}} + \frac{K_{ee}}{V_{ee}[S]}$$

Here the intercept is $\frac{1}{V}$ and the slope is $\frac{K_m}{V}$. There is one objection to this transformation, however, namely, that points representing very low substrate concentrations, which are likely to be maccurate, are given too great weight, since they become increasingly spread out in the plot of

 $\frac{1}{t_i}$ against $\frac{1}{[S]}$ A second type of transformation of the original Equation (5) is better in this respect. It is obtained by multiplying the last equation, (6), by [S] to give

(6a)
$$\frac{[S]}{t_1} = \frac{[S]}{V_m} + \frac{K_m}{V_m}$$

If $\frac{[S]}{l_*}$ is plotted against [S], the slope is $\frac{1}{V_m}$ and the intercept is $\frac{K_m}{V_m}$ From the intercept and the slope, Km is therefore readily obtained

^a Briggs and Haldane Biochem J 19 338 (19°o) Wilson Clapter in Lardy (ed.) Respiratory En ymee Vinneapolis Burgess Publishing Co. 01930 p. 17 ^a Linoweaver and Burk. J Am. Chm. Soc. 56 653 (1934) Respiratory Enzymes loc.

The affinity of enzyme for substrate is of the utmost importance and may eventually turn out to be the determining factor in deciding why an enzyme consisting of a protein plus a prosthetic group is so much more efficient as a catalyst than the prosthetic group alone

Synthetic Action of Enzymes. Enzymic reactions can be divided into

(1) Exergonic, or those which liberate energy

(2) Endergonic, or those which require addition of energy in order to occur

The reactions which result in the synthesis of glucose-1-phosphate from glucose, glutamine from glutamic acid and ammonia, acetylcholine, urca, peptides, riboflavin phosphate, coenzyme I, etc., are endergonic The energy for these reactions is supplied by the breakdown of adenosine triphosphate

Enzymic reactions can he divided also into

(a) Those which proceed to completion

(h) Those which proceed to a point of equilibrium at which both sub-

strate and products are present in measurable amounts

Examples of (a) are the decomposition of hydrogen peroxide by eatalase, the formation of rhodanate from evanide and thiosulfate by rhodanese and the synthesis of gum dextran from sucrose by dextran sucrase An example of (h) is the conversion of glucose-6-phosphate to glucose-1phosphate by phosphoglucomutase At equilibrium the digest will contain 5 per cent of glucose-1-phosphate and 95 per cent of glucoso-6phosphate.

One of the most striking demonstrations of the reversibility of enzyme action can be obtained with phosphorylase This enzyme which is found in animal tissues as well as tissues of plants, eatalyzes the breakdown of

glycogen or starch to glucose-1-phosphate

starch + H2PO, = glucose-1-phosphate

The glucose-1-phosphate (Corr ester) can be isolated readily from the system If glucose-1-phosphate is added to a potato phosphorylase preparation, starch amylose is rapidly formed and can be detected by the color it gives with iodine solution. For this reaction to occur it is necessary to prime the digest by adding a trace of glycogen or starch. Muscle phosphorylase likewise produces starch from glucose-1-phosphate Phosphorylases prepared from liver, brain, heart or yeast form glycogen from glucose-1-phosphate. These phosphorylases, ordinary phosphorylase, which forms 1-4 linkages, and the Q enzymes, which forms 1-6 linkages. The latter is not a true phosphorylase, but a transglucosidase.

The possibility of reversible or synthetic action of enzymes has thus been demonstrated, and it is of the greatest interest to know the part that such reversible enzyme action may play in the synthesis of proteins, fats, carbohydrates, and other substances in the animal body

OXIDATION AND REDUCTION SYSTEMS

Most if not all of the energy of living matter is derived from oxidative processes. Oxidation involves the loss of negative electrons from the sub-

stance oxidized, these electrons passing to the oxidizing agent which is simultaneously reduced. Thus the reaction of ferric chloride with potassium iodide involves a transfer of electrons from the iodide ions to the ferric ions so that the iodide is said to be oxidized and the iron reduced Biological oxidations involve the same transfer of electrons.

$$Fe^{+++} + 3Cl^{-} + K^{+} + I^{-} \rightleftharpoons Fe^{++} + 3Cl^{-} + K^{+} + I$$

In many oxidations the products of the reaction possess less energy than the reacting substances, and in these cases energy is liberated as heat or in some other form

Many oxidations are coupled with the formation of esters of phosphoric acid such as adenosinetriphosphate. Iuorganie phosphate is used to form these esters. The esters which are formed primarily are very labile, and there is a considerable free-energy decrease (ahout 10,000 calories) when they are hydrolyzed. Such an ester is adenosinetriphosphate, which contains two such labile phosphoric acid groups. These labile phosphoric acid residues are connected in a manner similar to that found in pyrophosphone acid and the hond is sometimes called a high energy phosphate hond. The formation of such compounds is a matter of considerable significance. It suggests how a cell might make use of the energy of oxidation For example, in the presence of hevokinase from yeast or muscle, the following reaction occurs.

 $adenosmetriphosphate \ + \ glucose \ \rightarrow adeuosmednphosphate$

+ glucose-6-phosphate

Thus glucose is chauged into a form in which it can readily enter into the intermediate reactions of carbohydrate metabolism. One might ask why phosphatase could not do the same thing with glucose. Phosphatase catalyzes the following reaction.

It is commonly considered that all enzyme reactions are reversible. However, thermodynamic considerations suggest that, although this reaction may be reversible, the equilibrium would be so far to the right that the may be reversible, the equilibrium would be so far to the right that the reverse reaction would be physiologically insignificant. Thus thero would be very little glucose-6-phosphate formed. On the other hand, in the reaction cited with adenosinetriphosphate, the equilibrium position is far to the side of glucose-6-phosphate and adenosinediphosphate. In this way it is thought that the cell can use at least part of the energy produced by exidation reactions to form materials which are useful to the cell.

Oxidation of Hydrogen. The first steps in the oxidation of fats, earbohydrates and proteins are dehydrogenations, brought about hy enzymes known as dehydrogenases. About 40 dehydrogenases are known In many instances the fats, carbohydrates, and proteins are not debydrogenated as such, but are first broken down by hydrolysis or desmolysis to genated as such, but are first broken down by hydrolysis or desmolysis to simpler compounds, ie diphosphogly cerie aldehyde, glycerophosphate, choline, amino acids, etc.

The individual dehydrogenases are frequently composed of a specific protein loosely united with coenzyme I or coenzyme II Coenzyme I is

also known as diphosphopyridine nucleotide, or DPN Coenzyme II is triphosphopyridine nucleotide or TPN

The hydrogen that is split off unites with these coenzymes, forming DP\H and TP\H The hydrogen may be next taken up by TP\-evtochrome C reductase, or hy DP\ cytochrome C reductase

TPNH + TPN-cytochrome C reductase

TPN

+ reduced 1PN cytochrome C reductase

This enzyme next reacts with cytochrome C

reduced TP\ cytochrome C reductase + cytochrome C ⇒ FPN-cytochrome C reductase + reduced cytochrome C + H*

Finally in the presence of atmospheric oxygen and cytochrome oxidase, the oxidation is completed

reduced cytochrome C + 211+ + 1/2O2 = cytochrome C + H.O

Some of the hydrogen present in substrates enters the Krebs cycle and is split off at one stage or another. One of these reactions involves succinic acid, which is oxidized by the succinic oxidase system. Here the initial reaction is

succinic acid = furnaric acid + 211
(succinic dehydrogenase)

The route -raveled by the hydrogen is probably as follows

succime acid \rightarrow cytochrome B \rightarrow cytochrome C \rightarrow cytochrome $\Lambda \rightarrow$ cytochrome oxidase

If some easily reduced dye, such as methylene blue is present it will be reduced to leucomethylene blue in the presence of succinic acid

succinic acid + Mb = fumaric acid + MbH2
(succinic dehydrogenase)

When leucomothylene hlue comes in contact with oxygen it is reoxidized, forming methylene hlue and hydrogen peroxide

$$MbH_2 + O_2 = Mh + H_2O_2$$

A number of yellow enzymes (old yellow enzyme xanthine oxidase, glucose oxidase of molds p-ammo acid oxidase) in the reduced state can react with gaseous oxygen to form hydrogen peroxide. In living cells this by drogen peroxide is very rapidly decomposed by the enzyme catalase. This reaction is

$$2H_2O_2 \rightarrow 2H_2O + O_2$$

Catalase is a heme-protein found to all living cells with the exception of certain hacteria. Its function appears to be the removal of the hydrogen peroxide which is toxic. Keilin and Hartree have observed that catalase can utilize hydrogen peroxide to oxidize methyl and ethyl alcohols to the corresponding aldehydes. On account of this reaction Theorell has sug-

gested that catalase is an important peroxidizing enzyme. However, this

hypothesis has little to support it at the present time

The hydrogen perovide may also be removed by peroxidase This enzyme, in the presence of H_2O_2 , oxidizes various phenohe compounds and amines, such as pyrogallol, guanaeol, hydroquinone, tyrosine, adrenatine, etc Peroxidase is a heme-protein, in fact, even hemin itself has a very slight peroxidase action Peroxidase, like other heme containing enzymes, is inhibited by HCN, H-S, and sodium azide

Oxidation by activation and transfer of hydrogen also occurs in the

oxidation of aldehydes First of all the aldehyde is hydrated

$$\begin{array}{ccc} R-CHO \,+\, H_2O \to R-CH-OH \\ & OH \\ & Aldehyde & Hydrated aldehyde \end{array}$$

The hydrated aldehyde is then oudized

This type of oxidation occurs in the oxidation of acetaldehyde to acetic acid by aldehyde dehydrogenase Similarly alcohol dehydrogenase mediates the oxidation of alcohols to aldehydes, as, for example, vitamin

A to retinene

Cytochromes. By use of the spectroscope, Keilin was able to demonstrate in hving cells the presence of three closely related substances which he called cytochromes A, B, and C Cytochromes appear to he present in all living cells except a few hacteria. The cytochromes are heme-protein compounds. The absorption hands of the reduced cytochromes may be easily observed by looking at a yeast suspension with a spectroscope. If a stream of air is hubbled through the suspension, the cytochromes are oxidized and the bands disappear. In animal tissues the cytochromes commonly cuts predominantly in the oxidized form

Oxidation by Activation of Oxygen. Warburg found in living cells a heme-containing compound which has the property of activating oxygen and which he called the respiratory enzyme. The activity of the enzyme is dependent upon the iron present, since HCN, in small amounts, and earbon monovide inhibit the action, apparently through combination with the metal. The inhibition by carhon monovide can be obtained only in the dark. Warburg's respiratory enzyme is apparently identical with the enzyme formerly called indophenol oxidase and now known as extochrome oxidase.

Cytochrome oxidase, in the presence of molecular oxygen, oxidizes reduced cytochrome C, as has already been described

Cytochrome C is reduced by such substrates as hydroquinone and by the Nadi reagent, a mixture of directhyl p phenylenediamine and α naphthol. Here the reaction is

Dimethyl- a-Naphthol Indophenol p-phenylenediamine

Yellow Enzymes. TPN-cytochrome C reductase and DPN-cytochrome C reductase are yellow enzymes? These enzymes contain ribofiavin (vitamin B; or G) in their prosthetic groups The prosthetic group of cytochrome C reductase consists of riboflavin phosphate (scalloxazine-b-ribose-phosphate) which is sometimes called a mononucleotides as prosthetic groups. An example is the old yellow enzyme of Warburg Other yellow enzymes centain prosthetic groups composed of riboflavin-phosphate-phosphate-bribose-adenine. Such a compound is called a dinucleotide Examples of yellow enzymes which contain isoalloxazine-adenine dinucleotide prosthetic groups are diaphorase, the Haas yellow enzyme, xanthine oxidase, and b amino acid oxidase. These yellow enzymes function in hydrogen transport as outlined above I tis the prosthetic group of these enzymes which is oxidized and reduced

Copper Enzymes. There are various enzymes which contain copper These are all inhited by HCV in much the same way as the iron-containing enzymes cytochrome oxidase, catalase, and peroxidase One of these copper enzymes is tyrosinase. This enzyme is apparently identical with monophenoloxidase and polyphonoloxidase. This enzyme oxidizes various phenohic compounds such as phenol, catechol, cresols, tyrosine, pyrogallol, and dopa (3.4-dihydroxyphenylalanine). For example

Water, not hydrogen peroxide, is always formed as a result of oxidation by this enzyme, and gaseous oxygen is used as the hydrogen acceptor Tyrosinase is also inhibited by H2S and by CO The inhibition by caroon monorade is not influenced by light Tyrosinase is commonly found in plant tissues, hacteria, and fungi, it also occurs in animals Tyrosinase is very similar to lacease which is also a copper-containing enzyme Lacease is found in various plant tissues and differs from tyrosinase in that it does not oudize tyrosine or p eresol

Ascorbic acid ovidase is another plant copper containing enzyme. It catalyzes the olidation of Lascorbic acid (vitamin C) in the presence of

oxygen

$$\begin{array}{c|c} O & O & O \\ \hline C & C & C \\ C & C & C \\ \hline C & C & C \\ C & C & C \\ \hline C & C & C \\ \hline C & C & C \\ \hline C & C & C \\ \hline C & C & C \\ \hline C & C & C \\ \hline C & C & C \\ \hline C & C & C \\ \hline C & C & C \\ \hline C & C & C \\ \hline C & C & C \\ \hline C & C & C \\ \hline C & C & C \\ \hline C & C & C \\$$

The dehydroascorbic acid may be reduced by hydrogen sulfide

It has been proposed by various workers that one or more of these copper-containing enzymes function in plant tissues in much the same way as does cytochrome oxidase in animal tissues. The matter is as yet unsettled

Sulfhydryl Group in Oxidations. Practically all tissues give a purple color with sodium introprusside and ammonia. This indicates the presence of the sulfhydryl group R-SH Such a group is found in the amino acid cysteine HOOC CH(NH2) CH2SH Cysteine is not however found to any extent in the tissues as such but is found in appreciable amounts in the form of glutathione, a tripeptide of glyeine, cysteine, and glutamic acid, which has the following structure

On oxidation cysteme may be converted into the disulfide form or cystine $R-SH+HS-R+O\rightarrow R-S-S-R+H_2O$, and the cystine may be reduced to cysteine $R-S-S-R+2H\rightarrow 2R-SH$ Glutathione may also be ovidized to the disulfide form $2G-SH \rightarrow G-S-S-G$ this reaction also being reversible The widespread occurrence of glututhione and the amounts found in eells indicate its importance but its mode of action is not yet clear Glutathione has been synthesized both by methods of organie chemistry and by the use of enzymes

New Coenzymes. In recent years the list of coenzymes has increased considerably Thus, we have undine diphosphoglucose necessary for the conversion of galactose-1-phosphate to glucose-1-phosphate by the enryme known as phosphogalactose isomerise Countyme III, found in yeast, is required for the oxidation of L-cysteine sulfinite acid to cystere acid

A yet more interesting coenzyme is coenzyme 1, containing pantothenic acid (q v , Chapter 35), adein he icid, and increaptothanolamine Coenzyme A functions in the acitylation of aromatic amines by pigeon liver homogenates in the presence of acetie acid and ATP. Here the intermediate product is acetyl coenzyme \ Coenzyme \ A is required for the acetylation of choline by nervous trishe Aged extracts of autolyzed pigeon liver convert oxalacetie reid to citric acid if coenzyme \ A, ATP, magnesium ions and acetate are present. Coenzyme \ A is necessary for the catalytic transfer of acetyl-bound phosphite, as well as for the arsenolysis of acetyl phosphate by the phosphotransacetylase of Clostradium Huyteri. In this process coenzyme \ A aceepts an acetyl group from pyruvic acid and forms acetyl phosphate. The enzymo present is phosphotransacetylase, which occurs neither in animals nor in yeast. Bacterial phosphotransacetylase together with an enzyme from pigeon liver can employ synthetic acetyl phosphate to synthesize acetoacetic acid.

Coenzyme A is thought to be involved in the synthesis of phospholipides. Here ATP supplies the energy It was found that yeast extracts convert coenzyme A to a pyrophosphate in the presence of ATP. This compound can react with acetate to give rise to acetyl coenzyme A and

pyrophosphate

6-Thiocite acid (lipote acid, or 68-dithio-n-octanoic acid) has been isolated in crystalline form from hiver and from yeast. This substance is necessary for the growth of the protozoan Tetrahymena gelix and is able to replace acetate needed for the growth of Laclobacillus case: It appears that lipote acid is necessary for the oxidation and dismitiation of pyruvic acid by Streplococcus fecalis, and that the formation of acetyl coenzyme A and succinyl coenzyme A requires the presence of both thuranne pyrophosphate and lipote acid. Thiocite acid is discussed further in Chapter 35

EXPERIMENTS ON ENZYMES

ROLE OF IRON IN OXIDATIONS

1 Catalytic Decomposition of Hydrogen Peroxide by Metals Transfer 5 ml portions of hydrogen peroxide to each of 4 test tubes To one add a very small amount of finely divided metallic platinum, to the second a small amount of powdered magnetite (Fe₀O₃), to the third a similar amount of ordinary ferric oxide or hematite (Fe₀O₃), and to the fourth a small amount of magnetite and a few drops of 1 per cent NaCN (Poinon) hore any evolution of gas After

^{*}If it is deemed advisable by the matricior to give all the practical work upon enzymes at it is pon this it course additional experiments will be found to the chapters on digestion at it is post in it is course additional experiments will be found to the chapters on digestion paper in platine chloride solution and give in the critical weder prepared by dipping fifter that magnetite produces active oxygen even when heated to 330° C. although it then has product sets free molecular not active oxygen even when heated still further to \(\delta \text{of} \text{ C} \) but the latter decomposition (Fe-O₂) are it has when heated still further to \(\delta \text{of} \text{ C} \) but the latter decomposition of fercours.

a few minutes add a few drops of an alcoholic solution of gualac. This is oxidized to a blue color by active oxygen. Which tubes show this reaction? Cyanide paralyzes cell respiration, apparently by Inactivating the iron which is an essential part of the system.

EXPERIMENTS ON PLANT OXIDASES

1. Demonstration of Potato Oxidases. It is convenient to combine the study of potato oxidases with a study of the composition of the potato. This throws light on the value of the potato as a food. It also gives information as to the

composition of a typical vegetable cell.7

(a) PREPARATION OF POTATO EXTRACTS Wash and peel a medium-sized potato. Grate rapidly and transfer the gratings at once to a piece of cheesecioth which is suspended in a beaker containing about 200 ml. of distilled water, Work gently with the hand to get out as much of the starch as possible. Keep this extract (water extract No. 1). Make a second extraction using 200 ml. of distilled water (water extract No. 2). Make a third extraction, and if this does not contain an appreciable amount of starch, discard it. Work the puip until it is practically starch-free.

(b) Tests on Pulp Work a portion of the pulp very thoroughly with water until it is practically free from starch as indicated by the iodine test. Test for protein, using Millon's test, and for carbohydrate by the Mollsch test.

(c) Tests on Water Extract No 1 Pour off the supernatant liquid from the extract when the starch has settled out. Filter. Test for reducing sugars by Benedict's test. Test for protein. Boil 20 ml. of the extract and filter. Test the fitrate for morganic chlorides, suifates, and phosphates. Indicate in your notebook what information you have obtained as to the food value of the potato.

(d) SEPARATION OF STARCH Combine the starch obtained from extracts No. 1 and No. 2. Wash by decantation with distilled water. Drain off the water and turn the beaker upside down so that the starch will drain; otherwise moids

may develop.

(e) EXPERIMENTS ON POTATO OXIDASE. Into each of a series of 5 clean test tubes introduce 5 mi. of potato extract (extract No. 2 filtered). (if the extract is kept over, it must be preserved with toluene.) Introduce other reagents according to the following series: (1) Potato extract + 10 drops of 1 per cent phenol,* (2) potato extract + 10 drops of 1 per cent catechol, (3) potato extract + 10 drops of guaiac solution, (4) potato extract + 10 drops of pyrogaliol solution, and (5) potato extract + 5 drops of α -naphthol solution + 5 drops of p-phenylenediamine hydrochloride solution. This combination of a-naphthol and p-phenylenediamine ls known as the Nadi reagent.

Mix the contents of the tubes by shaking. Watch for any color chaoges. If necessary let stand until the next laboratory period (add toluene), and examine again. In this experiment the phonoi C.H.OH, catechol C.H.(OII):, and pyrogaliol CaH, (OH); are oxidized with the production of brown-colored compounds. The gualaconic acid in the gualac is oxidized to gualac blue. In the last tube we have the production of indopheool from the α -naphthol

and phenylenediamine under the influence of oxidase.

Two ovidases appear to be active in these tests. Monophenoloxidase (also called polyphenoloxidase and tyrosmase) is responsible for the

This experiment is based upon the laboratory directions of Dr. William H. Welker The phenol, catechol (pyrocatechol), p-pheny lenediamine hydrochloride, and pyrogallol are 1 per cent aqueous solutions. The a-naphthol is a 1 per cent solution in 95 per cent alcohol For the guaiac solution dissolve 0 5 g of gum guaiac in 30 ml of 95 per cent alcohol.

oxidation of the phenol to catechol, then to o quinone, and finally with condensation reactions to brown compounds of uncertain composition

$$OII \rightarrow OII \rightarrow OOI \rightarrow brown compounds$$

Polyphenoloxidase also acts upon catechol to form o-quinone and then the brown compounds just mentioned It also acts upon pyrogallol forming purpurogallin

Cytochrome oxidase in the presence of cytochrome C oxidizes the p-phenylenediamine and in the presence of α naphthol there is formed indephenol

$$\Pi_2 N \longrightarrow N \Pi_1 + \longrightarrow O \Pi \longrightarrow \Pi_2 N \longrightarrow N = \longrightarrow -O$$

Peroxidase in the presence of hydrogen peroxide also gives this reaction At least part of the color obtained in the above test is due to the action of peroxidase since the color develops more slowly if catalase is added to the potato oxidase preparation

The exidation of the guaraconic acid of the guarac to guarac blue is apparently due to the action of the e-quinones formed as indicated above, on the guarac, rather than any direct action of the enzymes on the guarac itself except in the case of the slight peroxidase action due to the presence of some peroxide Peroxidase appears to exclude the guarac directly

2 Experiments on Potato Peroxidase Prepare a series of 5 tubes containing 5 ml portions of potato extract and 10 drops of oxidase reagents as in the preceding experiment Prepare still another series, but use potato extract previously boiled for 5 minutes Then to each tube add 10 drops of 3 per cent hydrogen peroxide solution Note whether oxidation takes place more rapidly than in Exp 3 where no hydrogen peroxide is used is the peroxidase destroyed by builing?

The potato has a greater peroxidase than phenoloxidase activity, which accounts for the more rapid action in the presence of H₂O₂

3 Resistance of Oxidase and Peroxidase to Heat Into each of 3 test tubes lint: duce 5 mi of potato extract Put in a water bath at 70° for 10 minutes. To the first tube add 2 drops of 1 per cent catechol solution and then to each 10 drops of gualac solution, and to the third tube only, 10 drops of hydrogen peroxide solution. The appearance of a bine color in the presence but not in the absence of hydrogen peroxide indicates that peroxidase is not destroyed at 70° but that the phenol oxidase is not stable at this temperature llow would you prepare a solution containing peroxidase hut not oxidase? The

fact that the addition of catechol does not suffice to bring back the direct oxidase action indicates that it is not the catechol compounds but the oxidase that is destroyed at this temperature.

4. Role of Catechol Compounds in Oxidase System. Into each of 2 clean test tubes introduce 5 ml. of potato extract. To one, add 1 drop of 1 per cent catechol solution. Let stand for 5 minutes. Then to each tube add 10 drops of guaiac solution. Let stand and note any color change.

Most potatoes contain relatively little of the catechol compounds. The addition of catechol therefore bastens the oxidation of the guaiae, since the oxidation of the guaiae is secondary to the formation of o-quinones from catechol compounds, the o-quinones being capable of directly oxidizing guaiae.

- 5. Preparotion of Peroxidase. Scrapings of horseradish may be extracted with alcohol and dried. On extraction with water these dried scrapings give a peroxidase solution free from oxidase. By a more complicated process a preparation showing about 1,000 times the activity of the original material has been obtained.
- 6. Determination of Peroxidase. Into a 250-mf. flask introduce 100 mf. of a saturated solution of leucomalachite green, 2 ml. of 0.166 N sodium acetate saturated with toluene, 1 ml. of hydrogen peroxide, 3 and from 0.0025 to 0.05 unit of peroxidase in not more than 5 ml. of solution. All should be brought to 20° C. before mixing and kept at that temperature. Measure 10 ml. of N 114804 into a small flask, and add all at once at the end of 5 minutes to the digestion mixture to stop the action. Rinse the small flask, with 5 ml. of water. After 15 to 30 seconds, neutralize with a little more than an equivalent amount of NaOH solution. Shake vigorously to eliminate hubbles of CO₂. Compare with a standard malachite green solution containing 10 mg. per liter, and set at 5 or 10 mm. The standard is made up in 0.05N acetic acid and preserved with toluene. Yellow artificial light is better than dayight for the comparison. Under the given conditions one unit of peroxidase forms 53 mg. of malachite eigen.
- Demonstrotion of Cotolose. Mix about 1 g, ground liver with 3 ml, water and add a few ml, 3 per cent hydrogen peroxide. Test the gas evolved for oxygen, using a elowing match.

EXPERIMENTS ON ANIMAL OXIDASES

1. Schordinger Reoction. Place 5 ml. of milk in each of 3 test tubes. Heat one to boilling and cool. To each tube add 1 ml. of 0.02 per cent methylene blue solution. To tubes 1 and 2 add 1 ml. of 0.4 per cent formaldehyde solution. Mlk by gentle rotation, add 1 to 2 ml. of paraffin oil, and put in a water bath at about 40° C. The milk in tube 2 should gradually decolorize. The

^{&#}x27;The dye should be recrystallized twice from alcohol, once from petroleum ether, and again from alcohol Saturate utraited 0.0 SN acute early with the dye at 20° C The solution freed from air under a vacuum will keep for months. If precupitation occurs, warm to reclusive,

¹⁸ Dilute 30 per cent hydrogen peroxide (reagent grade) 100 times. Titrate an aliquot with M104 after adding sulfuric acid and ddute further so that each ml. contains 0.25 mg. H104.

reaction is an example of anaerobic oxidation in the presence of a hydrogen acceptor (methylene blue) and may be written thus.

$$CH_{2}O + H_{2}O \rightarrow CH_{2}(OH)_{2} + M \rightarrow HCOOH + M·H_{2}$$

This reaction is given more slowly by milk which has been heated and more rapidly by milk which has a high bacterial count.

2. Study of Tissue Oxidations by the Methylene Blue Method of Thunbert and Ahlgren. Principle. Incly divided issue is suspended in a solution containing methylene blue, phosphate solution to regulate the aedity, and the substance whose action it is desired to determine The tube is evacuated, placed in a water bath, and the time required for the methylene blue to be decolorized is determined. This is a measure of the rate of oxidation in the mixture. The nature of the substances capable of being oxidized by the tissue can be determined and the influences governing the oxidation process studied.

Procedure. The tubell most convenient for use in this experiment is illustrated in Fig. 77, Into each of 3 tubes introduce 9.9 ml, of a mixture of 8 ml, of methylene blue 1 2000 and 6 ml, of a phosphate buffer of plf 7. Then add 9.1 ml, of water coths force tube of 6 1 M



UUM TURE (THUNBERG)

metro exert or 3 tubes introduce 9.9 ml, of a mixture of 8 ml, of methylene blue 1 2000 and 6 ml. of a phosphate buffer of pH 7. Then add 0.1 ml, of water to the first cube, of 0.1 M potassium succinate to the second, and of 0.1 M potassium sycrophosphate to the third. Then add 0.2 g, of finely divided washed rabbit muscleit to each. Evacuate each tube for 2 or 3 minutes with a strong water-jet filter pump. 18 Place the tubes in a constant-temperature water bath at 35° C. Observe from time to time and note when each tube just loses its last trace of blue color. Record the number of minutes required in each case. Do the succinate and glycerophosphate solutions decolorize first? What does this indicate as to the oxidation of these substances by muscle tissue?

flocculent and amorphous, this compound has the normal copper content (17 per cent) and gives crystalline glutathione on decomposition

Wash the cuprous salt with 0 5 N II,800, on the centrifuge and then with water until the washings are free from sulfate. If a centrifuge is used, shout 10 washings are required. Or use a device prepared with two Jean sintered glass filters (grade 4) of suitable size. Put the cuprous salt in one and place the other upside down on top, a water-tight joint being made with a piece of wide rubber tuhing. Connect the lower funnel to a reservoir of distilled water about a meter above it, and attach the upper one to a flask and filter pump. Wash until the filtrate remains free from sulfate after shaking the filtration apparatus.

Suspend the precipitate in four to five times its bulk of distilled water, and decompose with well-washed Its Filter Free from Its by a stream of hydrogen Evaporate to a small bulk in a vacuum desiccator at room temperature if the volume of filtrate is too large for this and distillation in vacuum is necessary, use a good pump and distil at 25° C If the mixture does not crystallize spontaneously before reaching a sirupy consistency, rub with a glass rod to start crystallization Yield, shout 15g For the crystalline form of glutathione, see Fig 78

PREPARATION AND PURIFICATION OF ENZYMES¹⁵

Purified enzyme preparations may be made from digestive secretions containing the enzyme. More often, however, the source is an animal or plant tissue. To obtain the enzyme in concentrated form it must be freed as far as possible from the mixture of substances making up the cell

In the case of animal tissues the material should be immediately refrigerated and used as soon as possible. It may be cut up with a sussors or run through a meat-chopper and then rubbed in a mortar with quarts and or ground glass. Freezing with liquid air or carbon dioxide will often aid comminution and check fermentative processes. The material may also in some cases be dehydrated by drying in a current of warm air or by dehydrating with acetone or alcohol followed by treatment with ether to remore fat, and then grinding very fine in a ball mill.

The comminuted tissue may then be extracted by use of a auitable solvent (e.g., water, salt solutions glycerol or solutions of definite pH) which will extract the enzyme with as little other material as possible. The mixture may then be filtered or centrifuged. Since enzymes are proteins, the greatest difficulties in enzyme purification are connected with the removal of protein contaminations. Methods used in the purification of proteins are therefore useful (see Chapter 6).

Just as with proteins, the exact procedure will vary in each case The procedures generally required include dialysis, precipitation by means of salts or by liquids such as alcohol or acctone, and selective adsorption and elution Dialysis, employing most commonly the collodion bag, removes soluble diffuelble substances including the eatts that may be used in precipitation Electrodialysis may also be employed By means of fractional precipitation with salts such as ammonium sulfate, separation from many protein ma terials may be accomplished in some cases alcohol may inactivate the

¹⁸ This subject has been reviewed by Schwimmer and Pardee Advances in Enzymol 14, 375 (1953)

enzyme Among the most useful adsorhents are different types of aluminum hydroxide For example, by choosing a sultable form of this adsorbent and the proper acidity of solution, separation of the enzy mes of the pancreas has been accomplished Associated with adsorption methods is of course the elution of the adsorbed material from the aluminum hydroxide-enzyme complex by suitable solvents, among which are solutions of alkalf phosphates of different reactions or dilute ammona. For the preparation of the different forms of aluminum hydroxide see footnote 16 Since preparations of pure

Alumina B Precipitate as for Alumina A After the precipitation stir for nct more than

half an hour Then dilute at once in 12 liters and wash by decantation as above

All winso Co Ammonia alum is used. Ammonia used should be standardized by titration and abould be measured accurately. Disolic 22 g ammonium sulfare in 600 in 10 of water Warm to 63° C and add 100 ml of 10 per cent ammonia. Warm quickly to 58° Add with Approvia mechanical stirring all at once a solution of 76° g of ammonia ultim in 150 ml of water this solution being previously warned to 55°. The temperature rises to 61° Do not let the temperature to below 58° C Ten monites after beginning precipitation separatia as rapidly as possible from the mother houor by centrifuging Transfer with 1500 ml of water containing 1.25 g of NH; to a 1500-ml flast, and shake Centrifuge Repeat using 25 g of NH. Each treatment with ammonia requires about 17 minutes. Then wash three times more using water only. The last superstants fluid should cream tirtled The whola procedure from the first precipitation should require about 2½ hours and must be expedited as the precipitation should require about 2½ hours and must be expedited as the precipitation of the contraction is made to the procedure from the first precipitation should require about 2½ hours and must be expedited.

Alumina Cβ The α compound changes into the β form on standing a few hours. The flocculent suspension becomes a plastic gel less soluble in acid. The β form changes gradually

(10 days to several months) into the \(\gamma \) form

Alumina C γ Precipitate as for the α farm Star for 15 minutes at 60° Transfer with 5 liters are more of water to a tail jar and wash by decentation. To the fourth, wish water add 80 ml of 20 per cost NH₃ to decompose the base sulfate. Wash 12 to 20 times more with water or twice after the supernatant fluid and longer becomes clear 50mm months standing under water is required for complete conversion to the γ form to take place. The precipitate becomes more flocutient and is smoduled in cold dilute or fairly strong HCl

ddmina D Disolve 130 g of pure commercial aluminum histroxide with 140 g of KOH 180 per centi in 900 ml of hot water Dulute in one liter and filer Dulute to 10 liters Precipitate by running a gentle stream if earboo doxide through the solution for two days Decant and wash 12 times with water contaming carboo doxide and then with distilled

water The final washings remaio turbid

Aluminum Metahydroxide If one of the above forms of aluminum hydroxide be heated suddenly with ammonia in a sealed tube in 250°C and kept at that temperature for 8 to 9

hours there is a complete conversion to the metabydroxide form

Kaolin Kaolin is best treated with acid before use To 500 g of kaolin add 1500 ml of pure HCl (sp gr 1 18) and warm very slowly so that the mixture begans to boil at the end of the first day and then boil for another day. Wash with water by decantation Repeat the treatment with acid and washing three times more. Wash with cold water until the washings slow practically no acidit but the kaolin itself reacts atromly acid on I timus pathers.

show practically no aeduty but the kaolin itself reacts atrongly acid on 1 timus paper Other Adsorbents Ferric by doxide magnesium oxide and stannic and silicid acids are used Substrates may also be used as specific adsorbents as tristearin for lipase casein for

trypsin and congulated egg albumin for pepsin

Elition Solutions of ammonia or of d sod um phosphate are most generally useful in

freeing adsorbed enzymes from combinations with alumina

Different ents mes are differently adsorbed. The properties of an ents me as far as advorption is concerned may change during purification. In a general way the more finely dispersed gels such as alumina A B and C have more adsorptive power that those with less surface such as the microcrystalline D and metal 3 droude. For numerous applications see Will stitler et al. Untersuchingen after Engine Berlin Julius Springer 1978. Also Grassmann Ergebniuse Physiol. 27. 40° (1928). Oppenheimer. Die Fermente und Ihre. Wurkungen, Leiping Theme Vol. 3 ist ded. 1929.

it Preparation of Adsorbents Alumina A Warm 250 g of Ali(SO₄): 18H;O in 750 ml of water to 55° C and pour the solutino all at once with the most vigorous mechanical stirring into 2 5 liters of 15 per cent amnoma warmed in 55°. The temperature rises to 55° and is kept between 55° and 60° C with cootinued stirring for half an hour Transfer to a 5-liter flask with a reflux condenser and boil gently for 48 hours. Dilute in a jar with water to 12 liters. Wash 3 times by decantation. Stir the precipitate with 500 ml of 15 per cent am monia to decompose the traces of base sulflate. Wash until for three successive times the wash water no longer comes away clear and the precipitate becomes a plastic gel from which the wash water can be completely poured.

5

crystalline enzymes are of the greatest interest, there follows a description of methods used in the preparation of certain of these. Methods for the preparation of active preparations of vegetable thoase and sucrase are also included since these are more readly obtained from regetable sources. Other enzyme preparations are discussed in the chapters dealing with digestion.

- 1. Preparation of Crystalline Urease.12 Dilute 158 ml. of redistilled acetone19 to 500 ml, at 22° C. (= 31.6 per cent acetone). Pour over 100 g. of Arico jack bean meal in a beaker. Stir 3 to 4 minutes and filter through a Schieicher and Schull, No. 595, or Whatman, No. 1, filter. Allow about 150 mi. to filter at room temperature. Complete the filtration in the ice chest at 2° to 2.5° C., over night, Centriluge off the crystals that form, using cold centrifuge tubes. Examine microscopicaliy (see Fig. 75, p. 308). Drain and stir up with 5 to 10 ml. of cold 31.6 per cent acetone. Centriluge again. Dissolve the crystals in 15 to 40 ml, of distilled water at room temperature, and free the solution from insoluble matter by centrifuging. The activity of the concentrated solution is not lost very rapidly provided the material is kept in the ice chest.
 - 2. Preparation of Crystalline Pepsin. Dissolve 500 g. of Parke Davis pepsin (U.S.P. 1:10,000) in 500 ml, of water and add 500 ml, of N 1f:SO. Add with stirring 1000 ml. of saturated VgSO, solution. Fliter through fluted paper (S. and S. No. 145014) and then with suction. Discard the filtrate. The precipitate must not stand at room temperature more than 24 hours.

PRECIPITATE 1. Wash twice with an equal volume of 34 saturated MgSO.

Filter with suction, Discard the filtrate.

PRECEPTRATE 2 Stir with water to a thick paste and run in M/2 NaOil until complete solution. Great care must be taken to avoid local excess of NaOll. The pli must never rise above 5.0. Add M/2 II:SO, with stirring until a heavy precipitate forms (pli about 3.0). Let stand 3 to 6 hours at 8° C. Fliter with suction.

PRECIPITATE 3 Stir with water to a thick paste at 45° C. Add M/2 NaOtt caretully until the precipitate dissolves (filter it cloudy and discard the precipitate). Place the beaker containing the precipitate in a vessel containing about 4 liters of water at 45° C., inoculate with pepsin crystals, and allow to cool slowly. Cooling should require 3 to 4 hours and a heavy crystalline precipitate should form at about 30° to 35° C, (see Fig. 76, p. 308). Keep the solution at 20° C. for 24 hours. Filter off the thick crystalline paste with suction.

PRECIPITATE 4 Wash with a small amount of cold water and then with 1/2 saturated MgSO, and keep under saturated MgSO, at 5° C. The filtrate may be treated with M/2 H2SO, to attain a pli of 3.0, the amorphous precipitate filtered off and treated like Precipitate 3.

3. Preparation of Crystalline Trypsin and Chymotrypsin. Remove the Pancreas from cattle within one hour after slaughter, and Immerse in cold N/4 II:SO. Drain off the acid, mince, and suspend for 24 hours in 2 volumes of N/4 II, SO, at 5° C. Strain through gauze. Add solid (NII,), SO, to 0.4 saturation. Filter. Saturate to 0.7 saturation. Let stand 2 days at 5° C. Dissolve in water and re-fractionate between 0 4 and 0.7 saturation. Dissolve the precipitate from 0.7 saturation in 0.25 saturated (NII,) 2SO4. Adjust to pli 5 0.

Sumner J Biol Chem 69, 435 (1926) 76, 97 (1926) Sumner and Hand J Biol Chem.
 149 (1928) For the recrystallization of urease see Dounce J Biol Chem. 140, 307 (1934) "Commercial acetone distilled over fused CaCl; and soda line, to remove the water and

Let stand 2 days at 25° C. Filter. Grystals are chymotrypsinogen. (Retain filtrate for trypsin.) Recrystallize chymotrypsinogen in 0.25 saturated (NH_d): SO_4 at pH 5 about 8 times. Dissolve in N/50 H, SO_4 . Adjust to pH 7.6. Add a trace of trypsin. Let stand 2 days at 5° C. Adjust to pH 4. Salt out in 0.7 saturated (NH_d): SO_4 . Filter. Dissolve in N/100 H: SO_4 . Let stand 24 hours at 25° C. Filter off chymotry psin crystals.

Take the filtrate from the chymotrypsinogen crystallization for the preparation of trypsin. Adjust to pli 4.0, precipitate in 0.7 saturated (NH₂),SO₄, and refractionate between 0.4 and 0.7 saturated (NH₂),SO₄. Filter. Wash the precipitate from 0.7 saturation with sat. MgSO₄. Dissolve in 0.4 M borate of pli 9.0. Cool to 5° C. Bring to 0.5 saturation with MgSO₄. Let stand 3 days at 5° C. Filter off the crystals of trypsinogen, Wash with 0.5 sat. MgSO₄. Dissolve in N/50 H₂SO₄. Bring to 0.5 sat. with MgSO₅. Add 0.4 M borate to give a pH of 9.0. Let stand 1 day at 5° C. and filter off the crystals of trypsin. Yield, about 6 g. from 7 liters of acid extract of the pancreas.

- 4. Preparation of Vegetable Lipase: Procedure of Willstinter and Waldschmidt-Leitz. Rub in a mortar to a paste 20 g. hulled castor beans. "Then with continued rubbing add 140 ml, of water in portions of 5 to 10 ml. Centrifuge for 15 minutes at 3,000 revolutions per minute. Three layers are formed. Pour off the upper creamy layer and retain. Pour off the clear water layer and discard. Rub up the residue with 140 ml. of water as before and centrifuge. Pour off the creamy top layer and combine with the first portion. Use this suspension for lipase experiments.
- 5. Preparation of Sucrase from Yeast. Introduce 100 g. of compressed yeast into a 400-mil. beaker. Warm the yeast to 30° C. by placing the beaker for some time in a water bath at a slightly higher temperature. Add 10 mil. of toluol. Stir thoroughly with a heavy glass rod. The yeast should become liquefied in 30 to 45 minutes. Add 200 mil. of water. Mix thoroughly and centrifuge. To the residue in the centrifuge tubes, add a small amount of water and mix well, then add more water at 30° C. to make a total of 200 mil. Stir and centrifuge. To the yeast residue after pouring off the water, add 100 mil. of water saturated with toluol and 10 mil. of toluol, and incubate at 30° C. over night. Dilute with 4 volumes of water. With vigorous attring, carefully add acetic acid (not over 1 N) until a test with methyl red indicates a pH of about 3.5 to 4.0. Centrifuge. To the supernatur fluid add some infusorial earth and filter. Neutralize with ammonia to complete change of color with bromocresol purple. Keep in a refrigerator.
 - 6. Preparation of Crystalline Catalase. Put fresh beef liver through a meat grinder four times. Extract the catalase at room temperature by the occasional shaking of each kilogram of ground liver for 10 min. with 11. of distilled water. To each liter of material add 480 mi. of chilided solution of chiloroform, 1 part; and ethyl alcohol, 2 parts. Shake violently for about 30 sec. and at once filter in the ice chest through Whatman No. 12 filter papers. The next day the filtrates will contain a sediment of catalase crystals. This sediment is centrifuged down in a refrigerated centrifuge and the supernatant liquid is discarded. The crystals are dissolved at room temperature in distilled water (30-40 mil. for each kilogram of liver used). The solution is

¹¹ The greatest care must be taken to avoid possoning by the ricin which is present in the meats

¹⁰ Mosimann' Arch, Biochem Biophys , 33, 483 (1951).

centrifuged to remove a white amorphous impurity, decanted, and chilled in the ice chest for 12 to 24 hr. Catalase will precipitate as needles.

QUANTITATIVE ESTIMATION OF ENZYME ACTION

The amount of enzyme present in any mixture is expressed in terms of its activity as compared with an arbitrary standard. Either the time required for a given amount of enzyme preparation to bring about a definite degree of conversion of the substrate or the amount of preparation needed to bring ahout a definite degree of conversion in a specified time may be made the basis of comparison. Inasmuch as enzyme action is greatly influenced by the pII of the solution, by the presence of inorganic salts and activators, and by temperature, it is important that the conditions in the digestion mixtures should be made as nearly identical as possible and that the enzyme if in inactive form should be properly activated.

Chemical reactions follow the law of mass action—that is, their speed is proportional to the product of the concentrations of the reacting substances. In most enzyme reactions the speed of decomposition of the substrate tends at first to be directly proportional to the amount of enzyme present, particularly if the substrate is present in large excess. Later the reaction may slow up due to decrease in the amount of substrate or combination of the enzyme with substrate or products of the reaction. In quantitative studies of enzyme action it is hest to have a large excess of substrate and to keep the time of the experiments within the period when decomposition is proportional to time. To establish these limits an experiment must be conducted and a curve plotted showing the extent of decomposition at different time intervals. In certain cases, direct proportionality does not exist even in the early stages. In such cases a curve prepared as above will suggest the rule to be applied in the calculation of results.

DEMONSTRATION OF THE REVERSIBLE ACTION OF AN ENZYME

The enzyme phosphorylase is found in plant and animal tissues This enzyme catalyzes the following reaction

glycogen (or starch) + morganic phosphate \rightleftharpoons glucose-1-phosphate

The reaction is reversible and either glycogen or starch can be used to form glucose-1-phosphate (Con ester) Whether glycogen or starch is formed from glucose-1-phosphate depends upon the source of the enzyme. Muscle phosphorylase (in rifro) and the phosphorylase from higher plants form a poly-saccharide which is probably identical with the amylose fraction of starch Yeast phosphorylase and the phosphorylase from higher brain, and heart form glycogen from glucose-1-phosphate. In this case a second enzyme, the Q enzyme, is present in addition to phosphorylase. This Q enzyme forms 1-6 inhages

The position of the equilibrium is influenced by pH. At pH 7 an equilibrium is obtained when the total phosphate is 23 per cent as Cori ester

and 77 per cent as morganic phosphate

1. Formotion of Glucosc-1-phosphate (Hanes method as modified by Sumner and Somers). Boil 8 g. of soluble starch with about 100 ml. of water. Cool and add 12 g. of Na;HPO4 and 5 g. of KHrPO4 dissolved in ahout 300 ml. of water. Add 100 ml. of potato-cyanide extract. Dilute to 1 liter, add toluene, and mix. Keep at 20° to 25° C. for ahout 24 hours.

Inactivate the phosphorylase by adding 0.1 N iodine until the solution gives a permanent reddish-brown color. Remove the lodine hy adding 0.1 N thiosulfate until all brown color has disappeared. Add 10 to 20 ml. of 2 per cent pancreatin and allow to digest until no more dextrin remains, as shown by the iodine test. (This will take three or four hours.) Add 40 g. of barium acetate and about 8 ml. of 28 per cent ammonia, or enough to make the suspension definitely alkaline to phenol red. Mix well, centrifuge, and nour the supernatant through cotton. To each volume of supernatant add 2 volumes of 95 per cent alcohol and centrifuge down the harium salt of glucose-1phosphate. Discard the supernatant. Stir the precipitate with 30 to 60 ml. of water and enough 2 N sulfurle acid to give a pink color with thymol blue paper. Now add saturated potassium hydroxide cautiously until the suspension just fails to give a hine or brown color with Congo red paper. Add 6 g. of trichloroacetic acid and mix. To every volume of the suspension add 2 volumes of 95 per cent alcohol and stir. Centrifuge down the suspended matter. Decant the clear supernatant solution. Add saturated potassium hydroxide to it until it is decidedly alkaline to phenol red. The di-notassium salt of clucose-I-phosphate usually separates as an oil. Chill overnight at 0° to 5° C. Next day filter off the crystais, wash several times with 95 per cent alcohol and then with acetone, and dry at 50° C. The yield will be about 3.5 g., and the product will be about 85 per cent pure.

The potato-cyanide is prepared by disintegrating about 325 g. of recently sliced potato in 100 mi. of 0.01 N neutralized potassium cyanide in a blender. The disintegrated mass is squeezed in cheesecloth and the juice is centrifuged to eliminate the starch.

The hexose-I-phosphate, or Cori ester, is C₁H₁₁O₁PK₂·2H₂O. Does a trace of it reduce Benedict's solution? Heat a trace of it in boiling water for 5 minutes with 1 ml. of N sulfuric acid. Neutralize with 0.1 N NaOH. Now test for elucose. Test also for inorganic phosphate.

2. Formation of Storch from Glucore-1-Phosphate. Prepare 5 to 10 ml. of 0.1 per cent glucose-1-phosphate in water. Add a drop of phenol red and enough 0.1 N hydrochloric acid to hring the alkaline solution to approximate neutrality. Place 1 ml. of the solution in a test tube and add about 1 ml. of potato-cyanide extract. Add a drop of 0.01 per cent boiled starch solution and mix. This primes the reaction. From time to time remove a drop or two of the digest and test on a porcelain spot plate thy adding a few drops of 0.01 N iodine solution. Is starch formed? What is the chemical reaction.

CELL RESPIRATION

The ultimate objective in the study of those enzymes which are found within cells is to apply this knowledge to an understanding of the metabolic processes upon which the cell depends for its maintenance and function. The contributions of enzyme chemistry to this subject have been numerous and important, but it is clear that the isolation of an enzyme or enzyme activity from cells does not necessarily give information as to the significance of the enzyme in the intracte processes of metabolism within the cell. An alternate method of approach, therefore, is to study

the metabolic behavior of the isolated intert cell or tissue, under as nearly physiological conditions as possible, and to integrate knowledge gained in this way with that acquired by the study of individual enzyme systems.

It was Warburg who first showed that animal tissues and organs (liver, kidney, brain, etc) could be prepared in the form of thin sections of minics which would continue to earry on metabolic processes (respiration substrate utilization) for many hours after removal from the number of the could be added to t



FIG 79 BARCROFT-WARBURG MANOM ETER WITH AT-TACHED VESSEL

body if placed under suitable conditions, and that such metabolism could be followed quantitatively. While it is obvious that cells under these circumstances are no longer under the control of nervous or hormonal mechanisms, metabolic data obtained by this method of approach do not disagree with results obtained on the intact animal where the two methods can be compared, and have furmished valuable information concerning the localization of specific metabolic processes in individual organs of the animal body, as well as the effect of different substrates, coenzymes, activators, inhibitors, etc., on cell metabolism

In the study of isolated cells and tissues, chief attention has been directed to the respiration, and specifically to the oxygen consumption. In fact, some have defined the respiration of cells in terms of oxygen consumption alone, but carbon dioxide production is an equally important phase of respiration, and any complete characterization of cell respiration must include both oxygen consumption and carbon dioxide production. In addition to respiratory data much valuable information has been obtained by the use of the Warburg tissue-slice technique concerning other metabolic characteristics of cells such as substrate utilization, fermentative (glycolytic) ability, glucoce and glycogen synthesis, urea formation, conjugation of foreign organic com-

pounds etc The discussion in these pages will be confined to respiration and glycolysis, two characteristics of cell metabolism which are readdy measured manometrically, but it cannot be too strongly emphasized that knowledge of this type is essentially incomplete until it is supplemented with precise information concerning substrate utilization and end product formation, this fact should be more generally recognized

Measurement of Oxygen Consumption. The basis of the Warburg method for the measurement of the oxygen consumption of living cells is the apparatus shown in Fig 79 This is sometimes referred to as the Warburg apparatus but more properly is called the Barcroft-Warburg apparatus since it was adapted by Warburg from that developed by Barcroft for the study of blood gases It consists of a suitable vessel, containing the maternal to be studied in the proper fluid medium, and attached in a closed system to a manometer for measuring changes in



gas pressure ²¹ Various types of vessels in common use are shown in Fig 80, many other types have been described for special purposes. In general, an ordinary Barcroft-Warburg vessel has a capacity of 15 to 20 ml, with a center well and one or more side bulbs, larger and smaller vessels have also been used. The type of bored side-bulb plug illustrated is much to be preferred over the more common solid plug.

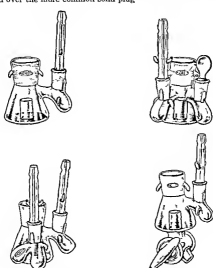


Fig 80 Types of Vessels Used for Manometric Studies on Cell Respiration

To provide for accurate temperature control, manometers and vessels are so constructed that the manometer may be mounted on the side of a constant-temperature water bath (thermostat), with the vessel completely immersed in the water. To ensure equilibration between fluid medium and gas phases in a vessel during an experiment, the manometer mounting is attached to vishaking device which shakes vessel and manometer horizontally at speeds which ordinarily amount to 110–115 oscilla-

n Complete equipment of the type described here (manometers glassware and thermostrium) be obtained from E Machlett and Son New York City Arthur II Thomas Co Philadelplus or American Instrument Co Silver Sterings Md.

tions per minute, with an excursion of 3-4 cm A complete assembly of

this type is illustrated in Fig. 81

The detailed application of the Warburg procedure for the measurement of oxygen consumption is given in the experiments which follow this discussion. The principle is as follows. The tissue, usually in the form of thin shees, is incubated at body temperature in a suitable buffered medium, in a vessel with attached manometer, the vessel containing oxygen rather than air. The center well of the vessel contains a little strong alkali solution which absorbs any CO₂ produced, so that any

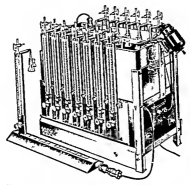


FIG 81 ASSEMBLY OF CONSTANT-TEMPERATURE BATH AND MANOMETERS AS USED FOR STUILES ON CELL RESPIRATION

pressure changes are due to oxygen consumption alone The side-bulbs provide for the addition of substrate activators, inhibitors etc., during an experiment if desired The gas phase in vessel and manometer capillaries is kept at constant volume, oxygen consumption is measured therefore by a fall in pressure, which is read on the manometer. The pressure readings when multiplied by a constant ("vessel constant") give the oxygen consumption, usually expressed in microliters (cubic millimeters). Readings are made at suitable intervals, they may be plotted graphically against time, or may be averaged to give the oxygen consumption over a given time interval. The standard method for expressing oxygin consumption is in terms of the number of microliters consumed in one hour by 1 mg (dry weight) of tissuic thus is symbolized by Qo-By convention the consumption of a gas is given a negative value Thus a Q_0 , of -60 represents the consumption of 6 microliters of oxygen per hour by 1 mg dry weight of tissue.

Representative Qo, values for various animal tissues are given in the table on p 340 These values must be considered approximations only. since it is known that the ovygen consumption of many tissues is affected by the presence or absence of various substrates, the previous nutritional history of the animal, the inorganic ion composition of the medium and the type of buffer used, the length of clapsed time between the death of the animal and the start of an experiment, and even the duration of the experiment itself, since oxygen consumption may not be linear with respect to time Despite these limitations, many valuable data have been obtained by this procedure, particularly where comparative and not absolute values are desired as in studies on the relative effects of various substrates, coenzymes, and inhibitors, comparison between normal and abnormal tissues, etc

A more serious limitation in the Warburg method, and one which is receiving increasing recognition, is that the tension of CO2 must be maintained at or near zero during the experiment, otherwise manometer readings will not reflect changes in oxygen content alone. This requirement means that metabolism is proceeding in the virtual absence of CO. and it is known that certain metabolic processes (e.g., the formation of urea by liver cells) require the presence of CO2, even the oxygen consumption of some tissues is significantly affected by the presence or absence of a physiological tension of CO2 Furthermore, measurements by the Warburg method are not possible in such media as normal blood serum or other bicarbonate-containing media, since such media require the maintenance of a finite tension of CO, to establish the pH "2 For the measurement of oxygen consumption and other respiratory characteristics of tissues in the presence of physiological tensions of CO2, the differential manometer described in the next section in connection with the measurement of CO2 production must be used

Measurement of Carbon Dioxide Production, Measurement of earbon dioxide production by respiring cells or tissues is technically more difficult than is measurement of oxygen consumption. There is no satis factory direct method for the continuous measurement of the carbon dioxide produced by cell respiration, as there is for the measurement of ovygen consumption. Hence relatively little work has been done on this phase of respiration This is unfortunate, for, as we now know, carbon dioxide is produced not by the direct oxidation of earbon compounds by oxygen but rather by decarboxylation of organic acids and in a manner essentially independent of oxygen consumption but presumably of equal importance to the cell Carbon dioxide production usually is expressed by giving the value of the respiratory quotient (R Q), which is the ratio of the volume of earbon dioxide evolved to the volume of oxygen consumed in the same time, hence if the Qo, and R Q are given, the carbon dioxide production is defined

One method for the determination of RQ in phosphate huffer using

[&]quot; For measuring oxygen consumption by the Warburg method in neutralized' serumi.e serum which i as been freed of bicarbonate by treatment with acid and evacuation—see Macleod and Rhoads Proc Soc Expli Biol Ved 41 268 (1939) Warren Am. J. Physiol 128 455 (1949)

the ordinary Barcroft Warburg manometers is to set up duplicate tissue preparations but to omit the alkali from the center well of one vessel Pressure changes in the vessel containing alkali are due to oxygen con sumption alone pressure changes in the second vessel without alkali represent the net (and opposing) effects of oxygen consumption and carbon dioxide production. At the end of the experiment by subtracting



FIG 82 CON STANT VOLUME DIFFERENTIAL MANOMETER.

the pressure change due to oxygen consumption alone as established on the first manometer, from the reading of the second manometer, the pressure change due to carhon droude production may be obtained Conditions in the two vessels including vessel constants must be as nearly alike as possible. This of course cannot be true with respect to the CO2 tension which is zero in the first vessel and continuously increasing in the second vessel. Results will therefore be in error if the CO2 tension influences metabolic processes within the cell and this is known to be the case with a number of different types of cells. This method is therefore not widely used. It is also possible to so arrange matters that only one

vessel containing alkali is used and the CO₂ absorbed by the alkali is measured by liberating with acid at the end of the experiment. This method is technically quite difficult is not too accurate and likewise suffers from the disadvantage that measurements are made in the

presence of a zero CO2 tension

To permit measurement of carbon dioxide production (and oxygen consumption as well) in the presence of a physiological tension of carbon dioxide and therefore in media such as blood serum or Ringer bicarbonate solution which are more nearly physiological than phosphateor similar buffers various types of so-called differential manometers have been developed One such type²¹ is illustrated in Fig. 82. It consists of two independent vessels and manometers with the manometers so arranged that pressure changes in the two vessels may be opposed to one another on the manometer to permit precase measurement of the pressure differences

each in use the two vessels are charged with identical amounts of medium and tissue the medium being either blood serum or Ringer bicarbonate and the gas phase being oxygen containing a physiological concentration (usually 5 per cent) of carbon droude At the beginning of the experimental period the tissue in one vessel is killed by tipping in acid from a side bulb and the tissue in the other vessel is then allowed to respire for the desired length of time after which it too is killed by the addition of acid Although the total pressure in both vessels is now quite high due to the liberation of CO₄ from the decomposed bicarbonate in the medium

²² Summerson J Biol Chem. 131 579 (1939)

by opposing one vessel pressure against the other on the manometer only the difference in pressure caused by the respiration of the tissue in the second vessel during the experimental period will register on the manometer This pressure difference is due to hotb ovygen consumption and carbon drovide production After it is noted, alkali is admitted into both vessels through a bottom stopcock (see Fig. 80 for the type of cssel used) and all the carbon dioxide is absorbed Again the differential reading is noted, this is now due solely to oxygen consumption by the respiring tissue. The carbon dioxide production is obtained by subtracting this reading from the first differential reading. Thus the oxygen consumption and R Q are accurately measurable under a continuous physiological tension of carbon dioxide. The only disadvantage of this method over the simple Warburg method as far as over consumption measurements are concerned is that the Warhurg method gives con tinuous readings, whereas the differential method gives but one reading, representing the entire experimental period Conditions therefore must he selected so that respiration is linear with respect to time, or nearly so

The use of the differential manometer is technically somewhat more difficult than is that of the simple Barcroft-Warhurg manometer, but it is the only accurate basis for the measurement of cell respiration under conditions which closely simulate those prevailing normally, and the increased significance of the results obtained more than justifies the

technical difficulties

A further advantage of the differential manometer is that it permits the measurement of aeroline and production (aeroline glycolysis see helow) by respiring tissues. If the tissue produces and—e.g., lactic acid—aerobically in a medium containing hearhonate neutralization of the acid results in the decomposition of an equivalent amount of hicarbonate to produce gaseous CO₂. This CO₂ is indistinguishable manometrically from the CO₂ of respiration unless the differential manometer is used, where it is readily done by a simple method which need not be described here (for details see Summerson (loc cit) and also hooks by Dixon and by Umbreit, et al., cited in the Bibliography). Since the production of acid aerobically is a characteristic which differentiates tumor tissues from many (but not all) normal tissues (see table on p. 340), the differential manometer is useful in metabolic studies on such tissues.

Anaerohic and Aerobic Glycolysis Practically all tissues show the ability to produce lactic acid from either glucose or glycogen under naerobic conditions. Such production of lactic acid is known as glycolysis although strictly speaking this term implies merely a disappearance of carboby drate rather than the specific production of lactic or other acids. With certain tissues—e.g., liver, kidney—the anaerobic formation of lactic acid is at the expense of tissue glycogen and is independent of the presence of glucose, it has been proposed that the term glycolysis be restricted to such origin of lactic acid from glycogen and that the term glucolysis be used where glicose is the source of the lactic acid.

Anaerobic glycolysis by tissues is readily measured with the simple Barcroft-Warburg manometers by suspending the tissue in a bicarbonate coataining medium in equilibrium with the proper tension of CO₂ in the

gas phase to maintain a suitable pH, but with introgen rather than oxygen present. Since in the absence of oxygen there is no respiration, pressure changes on the manometer will ordinarily be due solely to decomposition of bicarbonate by the acid produced anaerobically. Anaerobic glycolysis usually is represented by the symbol Q₂*, and is expressed in terms of microliters of CO₂ produced by the action of acid on the bicarbonate present per milligram dry tissue per hour. The anaerobic glycolytic power of various animal tissues is given in the table below. No distinction is made in this table between glucolysis and glycolysis, and the values for certain tissues such as liver and kidney are subject to considerable variation because the rate of glycolysis usually falls off continuously throughout the average experimental period. It will be noted that various tissues differ considerably in their Q₂* values and that tumor tissues in general (but not exclusively) are quite high in this respect.

RESPIRATION AND GLYCOLYSIS OF SELECTED ANIMAL TISSUES*

Tusaue	Qo,	RQ	$Q_G^{O_2}$	$Q_{\sigma}^{N_{\theta}}$
Muscle (dog)	2	0 90	0	4
Panereas (dog)	3		0	4
Bone marrow (rabbit)	5	0.90	2	13
Rous sarcoma (chicken)		0.93	20	30
I iver (fetal rat)	5 7	1 00	0.5	8
Testis (rat)	8	0 90	4	8
Jeasea sarcoma (rat)	1 9	0 78	17	34
Liver (adult rat)	1 10	0 5-1 0	0.5	3
Embryo (chicken)	11	1 00	2	18
Intestinal mucosa (rat)	12	0.85	2	4
Spleen (rat)	12	0 89	2	8
Brain (rat)	13	1 00	2 2 2 0	19
Thy road gland (rat)	13		0	2
Retina (rat)	19	1 00	22	88
Lidney (rat)	21	0 83	0	3
Chorion (rat)	26	1 00	7	32

^{*} Many values are averages from the I terature. For sign ficance of symbols used, and further discussion, see text.

The production of acid aerobically is also measurable in terms of the decomposition of bicarbonate is symbolized by $Q_0^{\rm co}$ and expressed in the same units as for anaerobic glycolysis Aerobic glycolysis is more difficult to measure than anaerobic glycolysis since there must be a way to differentiate the ${\rm CO_2}$ of acid production from the respiratory ${\rm CO_2}$. The only successful way to do this is with the differential manometer, as discussed previously. In contrast to acid production anaerobically, which in the majority of cases is quantitatively due to lactic acid, only a fraction of the acid produced aerobically is usually accountable for as lactic acid. Little specific information is available concerning the nature of other acids produced under these circumstances. As can be seen from the data above, most normal adult tissues have a relatively low aerobic glycolysis, certain specialized normal adult tissues and all tumor tissues, are cliaracterized by a significant aerobic elveolysis.

It is felt by some that the inhibition of anaerobic glycolysis ("fermentation") by aerobic metabolism, which is called the Pasteur effect, may be associated with the presence of a specific enzyme (Pasteur enzyme) whose function is to orient cell metabolism into either fermentative or oxidative pathways. Evidence for the existence of such an enzyme is limited at the present time

EXPERIMENTS ON CELL RESPIRATION

I Colibration of Vessel and Manameter Principle For the calculation of changes 10 gas content in a vessel from the manameter readings under a particular set of experimental conditions it is necessary to know the entire volume of gas space in the empty vessel manameter sude-arm and manameter capillary down to the level of manameter fluid as it will be placed in actual use. The best way to do this is by filling the entire space with mercury, then collecting and weighing the mercury, from the known density of the mercury at the temperature of use, the volume occupied by the mercury is readily obtained.

Procedure (a) Place the manometer (clean, dry, and empty of manometer fluid) in an upright position Fill the clean dry vessel with mercury, being sure that the vessel side tap is firmly in place (it may be lightly greased and held secure by springs or wound rubber bands). The mercury must fill the entire vessel including the center well and side bulb, any trapped air bubbles may be dislodged with a piece of fine wire. The mercury should come to within about one-half inch of the top rim of the vessel. Holding the vessel in the hand, set it slowly in place on the manometer side arm until it is firmly seated Excess mercury will run out between vessel and manometer groundglass surfaces during this process, should be brushed off if adhering to the glass, and may be caught in a large tray set beneath the entire assembly for this purpose A thread of mercury will likewise be forced up into the capillary of the manometer side arm for a variable distance. When the vessel is firmly seated on the manometer, make a mark on the manometer side arm to Indicate the limit of the mercury thread. The best position for this is in the straight portion of the side arm capillary a few centimeters above the vessel, other vessels may then be calibrated on this same manometer in the future, without the necessity of removing any manometer fluid present, by adjustment of the mercury thread to the same mark, this adjustment is particu larly easy if the vessel side built tap is bored as shown in Fig. 80, since by turning the tap to open it slightly, by slight pressure or suction the mercury level in the manometer side arm capillary may be controlled at will

When the mark has been made, slowly withdraw the vessel from the manometer, allowing the capillary thread of mercury to follow and be caught in the mercury in the vessel as it is removed from the manometer Transfer the mercury from the vessel to a tared beaker, and weigh the mercury to the nearest 0 1 g Note the temperature of the mercury From the volume occupied by 1 g of mercury at the indicated temperature, a calculate the

[&]quot;The volume occupied by 1 g of mercury at various temperatures is as follows

Temp °C	I olume mi	Temp °C	l olume ml
16	0 0 377	24	0 0 388
18 20	0.073*9	26 28	0 0"390
22	0 67395	30	0 0*396

volume occupied by the mercury at this stage of the calibration Call this volume V,

(b) Invert the manometer, turn it so that the front faces to the right and the aide arm projects to the left and fasten it accurely in an upright position with a large adjustable huret clamp attached to a ring stand Now loosen the attachment of the buret clamp to the ring stand and tilt the manometer clockwise until the calibration mark on the aide arm and the 150 mm graduation mark on the attached manometer limb are in line with each other and parallel to the table top Clamp the manometer firmly in this position Attach to the aide orifice of the manometer stopcock a piece of rubber tuhing carrying a screwcock and connected to a leveling hulb containing mercury The manometer stopcock should be lightly greased and may be held secure by rubher hands. Turn the manometer stopcock so that mercury may enter from the rubber tuhing and rise into the manometer capillaries By carefully releasing the screwcock on the rubber tubing, allow mercury to enter the manometer capillaries and rise in the aide arm and main limb until the mer cury level reaches both the calibration mark on the side arm and the 150 mm graduation on the main fimh Slight further tilting of the manometer may be necessary at this point If there are any trapped air hubbles lower the leveling hulb to retract the mercury and repeat the adjustment

When both capillarles of the manometer are filled with mercury to the desired points, turn the manometer stopcock very slightly to cut off the flow of mercury, place a tared beaker under the straight capillary orifice of the manometer stopcock, and then turn the stopcock so that the mercury drains from the manometer capillarles into the beaker. Any droplets of mercury remaining behind may be forced out and into the heaker by hlowing down the capillarles. Weigh the mercury, determine its temperature, and calculate the volume of the side arm and main limb capillarles. Call this volume $V_{\mathcal{U}}$

CALCULATION The total volume of empty vessel and manometer gas space 1 τ 15 equal to 1 γ + 1 ν from the value of 1 τ the vessel constant I under a particular set of experimental conditions is calculated as follows

$$k = \frac{1}{P_{\bullet}} \frac{273}{T} - \frac{1}{I_{\bullet}} \left(\frac{273}{T} - \alpha \right)$$

where I is the total solume of sessel and manometer as described above and I is the volume of fluid medium in the vessel during an experiment both of these volumes are expressed in microliters (cul is millimeters) rather than in millilaters (1 m) = 1000 microliters) P_i is the equivalent of I atmosphere pressure in min of manometer fluid (700 for mercury 10000 for Brode fluid see below). This the temperature in degree Absolute at which measurements are made and a is the absorption coefficient (solubility) of the gas undergoing absorption or evolution is For the derivation of this equation see the book by Dixon the form of the equation is that described by Macieod and Summerson is

It can be seen that the constant for a particular versel and manometer depends upon the zize of the vessel the volume of medium used the temperature and the nature of the gas concerned. The following example illustrates the calculation of the vessel constant for oxygen k_{0r} for a particular vessel and manometer where 1 τ = 15 70 mi and 20 mi of medium are employed at 38° C

¹³ At 35° C. α for O₂ is 0.024 for N₂ 0.012 and for CO₂ in water 0.55 in Pinger solution 0.377 and in Pinger solution containing 0.3 N HCl 0.517 For values of other gases and at other temperatures are bandbooks giving physical constants of gases 13 Macleod and Summerson. Science, 81, 220 (1940)

$$V_T = 15.75 \text{ ml} = 15,750 \text{ µl}$$

 $V_P = 2.00 \text{ ml} = 2.000 \text{ µl}$
 $T = 38^{\circ} \text{ C} = 311^{\circ} \text{ A}$
 $P_e = 10.000$
 $q_{D_e} = 0.024 \text{ nt} 38^{\circ} \text{ C}$

therefore

$$\begin{array}{l} \lambda_{Or} = \frac{15}{10} \frac{750}{0000} \times \frac{273}{311} - \frac{2000}{10000} \left(\frac{273}{311} - 0024 \right) \\ = 138 - 017 \\ = 121 \end{array}$$

The ressel constant will be different with other amounts of medium at other temperatures and for other gases. For routine work on oxygen consumption at 38° C, the only rariable is likely to be the volume of medium employed. Tessel constants should therefore be calculated for the rarious volumes of medium apt to be used or the simple graphical method described by Macleod and Summerson (loc cat.) for this purpose may be employed.

The cultivation is described for a manumeter fluid setting at the 150-mm mark on the graduated limb. This is the most satisfactory position but cultivation obviously can be made to some other setting. If a thread of mercury is placed in the graduated capillary, the length in mm noted and the mercury then weighed and its volume computed the volume of the graduated capillary per mm may be obtained and from this it is possible to calibrate the manometer at any setting on the scale provided the calibration at one setting is also known. This is of value under certain circumstances.

MINOMETER FLUID The most commonly used manometer fluid is the aqueous salt solution described by Brochett with a specific gravity of 1033, so that 10,000 mm are equal to I atmosphere (760 mm of mercury), this makes for obvious convenience in calculation, with a much greater sensitivity than mercury. For special purposes, however, any other fluid whose P_e value is known (water, mercury, paraffin oil, etc.) may be used

The manometer fluid usually is contained in a stoppered rubber tube attached to the manometer which acts as a reservoir, controlled by the pressure of a screwcock. The fluid may gradually leak out or be otherwise lost, it may be repleashed at any time, even during an experiment, by inserting a hypodermic needle attached to a syringe containing extra fluid through the rubber tube wall, at a slight angle to form a "Bursen valve" after withdrawal of the needle. It may also be added through the open top of the manometer, with precautions to avoid trapping air bubbles Should such bubbles be present, they may be forced to the top of the fluid column and dissipated by alternate pinching and release of the rubber tube reservoir with the fingers, to force the manometer fluid up and down.

2 Measurement of Oxygen Consumption of Rat Liver Slices by Warburg Method (a) Preferation of Tessue Kill a young adult rat by a blow on the

²¹ Brode fluid Is made as follows: Dissolve 23 g of sodium chloride and 5 g of bile alits codum tarurogly excholate or sodium ply excholate Merck) in 50 ml of water A few drops of an alcoholic thymol solution may be added as a preservative and it is convenent to color the fluid by ad ling a few hundred mg of a suitable dge (Crystal volect Gentian violet or Fv ans blue). The specific gravity of the final solution should be 1 033 i.e. 10 ml should weigh 1 033 times as much as 10 ml of water at the same temperature.

head, followed hy severing the neck vessels and spinal cord at the neck with scissors Open the andominal cavity and remove the liver as quickly as possible Rinse the liver in a heaker containing Ringer's solution," then cut off the largest lobe of the liver and place flat on a pad of filter paper to drain momentarily With a sharp thin razor blade moistened with Ringer's solution, cut off a small portion of the liver in such a way that the cut surface makes an angle of about 45° with the table top, and discard this portion Continue cutting freehand along the plane of the first cut surlace to obtain a number of thin slices of liver, making each slice as uniformly thin as possible (the leading edge of the razor blade should be visible through the slice as it is being cut) With practice, uniform silees about 0 5 mm thick are readily obtained As each slice is obtained, transfer it to a flat dish containing Ringer s solution, and keep the hlade of the razor and the eut surface of the liver moistened with Ringer's solution Continue cutting until sufficient silces for the experiment are obtained, usually two to lour slices are required per vessel

(b) PREPAINTION OF VESSELS Measure duplicate (or triplicate) 2-ml portions of medium" into clean dry Barcroit-Warburg vessels, avoiding the center well. In another vessel place a few ml of water, this will he the thermo harometer control With a lorceps or platinum wire, transfer a liver silce from the dish to a pad of filter paper and drain momentarily, then quickly transfer to a tared watch glass on a balance Add more silces treated in the same way until the desired weight of liver tissue is obtained, 100 to 200 mg wet weight is usually satislactory Immediately transfer the weighted clump of silces to one of the vessels, immersing them in the medium. The vessel may be shaken briefly to separate the silces in like manner, charge the remaining vessel scontaining medium with weighted amounts of tissue. The weights of tissue in each vessel need not be identical, but they should be known to the nearest milligram.

When all the vessels are ready, obtain one more known weight of tissue by exactly the same procedure, but transfer this to a small tared watch glass of weighing dish. This is to be dried in an oven overnight at 100° C, and weighed again, to determine the dry weight of the tissue

When the tlasue is in place in the vessels, complete the preparation of the vessels by placing in the center well of each (except the thermobarometer vessel) 0.2 ml of to per cent KOH solution, using a pipet with a fine the No alkali must get into the medium surrounding the center well. It is advantageous, but not absolutely necessary, at this point to insert a small roll of starch iree filter paper (Whatman No. 40 is satisfactory) in the center well so that the top of the roll projects a millimeter or two above the rim of the center well, the alkali, being absorbed on the paper roll, thus projects into the gas space of the vessel and is better able to absorb CO₂.

(c) PREPARATION OF MANOMETER When all the vessels are ready, mount each one on its manometer, using a small amount of suitable stopcock grease,

^{**} Runger's solution may be prepared as follows to 960 ml of 0 154 M NaCl solution add **O ml of 0 154 M KCl solution and 20 ml of 0 11 M CaCl; solution

¹⁰ The usual me hum is Binner pl coshists. To 10 volumes of Ringer a solution and I volumes of 1/15 pl copiate huffer pH 7.4 For 10 volumes of Ringer a solution and I volume of W1/15 pl copiate huffer pH 7.4 For paratison of pl cophate buffer see Chapter I if glucose is desired in the medium a level of 200 mg per 100 ml map, because of which add ng 2 ml of 10 per cent glucose solution to 10 ml of med um The presence of glucose in the med um does not affect the oxygen consumption of liver slaces but is necessary with many other trausses.

^{**} Cello-Seal obtainable from the Fisher Scientific Co New York is quite satisfactory Ordinary petroleum jelly or anhydrous tanolin may also be used but since these

and attaching the vessels firmly to the mannmeter shanks hy small springs or wound rubber bands. Fill each vessel containing tissue with oxygen by passing a slow stream of the washed gas from a tank. It through the vessel, attaching the rubber tube delivering the gas to the top of the manometer so that the gas passes through the manometer stopcock and capillaries into the vessel and out the opened side bulb of the vessel. A few minutes' passage is sufficient for each vessel; all the vessels can be gassed at once if a multiple manifold attached to the tops of all the manometers is used. It is not necessary to gas the thermobarometer vessel.

When the vessel is filled with oxygen, turn the stopcock at the top of the manometer so as to close off the manometer and divert the flow of gas to the outside air; at the same time, have the greased plug for the side bulb ready, and as soon as tha gas flow has been diverted, insert this plug and fasten it securely. Continue in this way with each of the other vessels being gassed.

Place the manometers one by one on the constant-temperature bath, which should be at 38° G. As each vessel enters the thermostat, the temperature rise will cause expansion of the enclosed gas and the manometer fluid will rise. Release the excess pressure by momentarily opening the manometer stopcock to the air and then closing it. This operation may have to be repeated. The thermobarometer is placed on the bath with the manometer stopcock open.

Start the shaking device and shake ressels and manometers for 10 minutes; this is usually sufficient to bring about temperature and pressure equilibrium. During this time, adjust the manometer fluid so that the level in the closed limb is approximately at the calibration setting (usually the iso-mm. polat on the scale). Since the pressure change to be expected during the experiment will be negative, the level of manometer fluid in the open limb should be as high as possible, to permit maximum range of readings. It may be necessary here also to release the pressure within a vessel momentarily to permit the indicated adjustment of manometer fluid level in each limb. During the preliminary equilibrium period, close the thermobarometer also; the level of manometer fluid should be about the same in each limb here.

When measurements are to be hegun, stop the shaking device, note the time, and immediately set the manometer fluid level in the inner (closed) limb of the first manometer at exactly 159 mm. Record the fluid level in the open limb, to the nearest mm. Now read the thermobarometer in the same way. For each vessel reading there must be a thermobarometer reading unless temperature and atmospheric pressure remain sufficiently constant so that one thermobarometer reading suffices for all the vessels; thus is rarely true.

Continue with the reading of the second manometer in the same way, until all of the manometers containing tissue have been read. When this has been done, start the shaking device, Repeat the readings as above, at suitable time intervais, usually 10 or 15 minutes, for the duration of the experimental period.

CALCULATION Subtract each reading for a vessel from the previous reading, and from this subtract algebraically the change in thermoharometer reading during the same time, to obtain the net change in pressure for the time interval covered by the readings Multiply the pressure change, h, by the vessel constant for oxygen under

become somewhat soft at 35° C, it usually is necessary to tighten the vessels on the manometers at least once during the preliminary incubation period when they are used

¹¹ Tanks containing oxygen and other gas mixtures mentioned here may be obtained from the Ohio Chemical Co., New York City

cedure above If at the end of an experimental period the fluid level in the open limb is below the graduated range when the manometer is set at 150 mm and it is desired to obtain a reading without resetting the manometer, set the manometer level at two successive points above the 150 mark—e.g., at 170 and at 160—make each reading quickly, and extrapolate to the reading at 150, since the change in reading between 160 and 150 will be the same as that between 170 and 160

3 Measurement of Anaerobic Glycolysis by Tissues Prepare the tissue as described under Exp 2, but use Ringer-blearhonate solution containing added glucose as the medium. and do not put any alkali into the center wells After attaching the vessels to the manometers, pass a stream of washed nitrogen gas containing 5 per cent carbon dioxide. through the vessels for 20 to 30 minutes at room temperature to completely displace any arygen present During the gassing period run the manometer fluid up and down once or twice to replace the air in the manometer capillary by the gas mixture, and shake the vessels occasionally to aid In displacing any air hubbles trapped in the medium. In turning the gas off and closing the vessels, take suitable precautions against the possible diffusion of room air into the system in the process. Vessels with hored plugs in the side hulhs, which can be left in place but clear for the passage of gas during the gassing period, and then closed by a quarter-turn, are much to be preferred over the type having solid side-hulh plugs.

The rise in pressure caused by immersing the vessels in the thermostat at 38° is released as described under Exp 2, again with precautions against allowing air to reënter the system. Since the readings will always be positive in this type of experiment (i.e., the pressure will increase owing to decom position of hicarbonate by the acid produced), set the manometer fluid level in the open limb at the bottom of the scale rather than at the top. This is easily done by forcing up the manometer fluid in the closed limb to about 200 mm, opening the stopcock cautiously to release most (but not all) of the excess pressure then closing the stopcock and lowering the manometer fluid level to the 150 mark, on the closed limb. If this is done properly, the fluid level in the open limb will be low down on the scale at this point.

Equilibrate by shaking for 10 minutes, then start readings, making them against a thermobarometer (which need not be gassed) just as described for the measurement of oxygen consumption in Exp 2 Read at sultable intervals for the desired length of time

CALCULATION Subtract each reading from the next one and subtract algebraically from this the change in thermobarometer reading during the same period as for Exp. 2, to obtain the net increase in pressure due to and production Multiply by the vessel constant for CO₂ at the fluid volume used $k_{\rm CO}$, to obtain the number of μ of CO₂ colved as the result of send production Express results in terms of $Q_{\rm C}^{\rm in}$, which is defined as the number of microliters of CO₂ equivalent to the acid produced by 1 mg dry weight of tissue in one hour, under anaerolic conditions

nt To 5 volumes of Ringer a solution add 1 volume of 0.15 M NaIICO; solution Bubble the nitrogen (05 per cent) CO₂ (5 per cent) gas markere through this solution for about 15 minutes before using to bring to pH 7.4 A glucose concentration of app roximately 200 mg per 100 ml may be established by adding 2 ml of 10 per cent glucose solution to each 100 ml of medium

³¹ Tanks containing 95 per cent nitrogen 5 per cent carbon dioxi le gas mixture may be obtained from the Olio Clemical Co. New York City. The gas mixture may be freed from the traces of oxygen usually present by passing through a heated tube filled with metallic copier turnings. Pass the dried gas through a wash bottle containing water to saturate with water yapor before entering the vessels.

With certain tissues anaerobic glycolysis proceeds linearly for hours, provided sufficient glucose is present, with other tissues the rate of glycolysis is independent of the presence of glucose (and hence is presumably at the expense of glycogen) and may show a continual decrease with time. In this latter instance, the Q values represent the average for an hour, or may be computed on the basis of shorter experimental periods.

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13

Salivary Digestion

Digestion in General. The greater part of the food elements in the diet require special treatment to render them capable of absorption and utilization by the body. Water, glucose, and eertain inorgauic salts and vitamins are exceptions, but the proteins, fats, and carhohydrates, as well as other substances, must be split up into simple components as monosaccharides, glycerol, fatty acids, amino acids, etc. The necessary changes are chiefly bydrolytic in character and involve especially tho action of enzymes found in the different parts of the gastrointestinal tract. Because of the varieties of the quantity and character of food ingested and the variety of changes that must be brought about, the different parts of the gastrointestinal tract must show a considerable power of adaptation and coordination. In this, both bormone and nervous mechanisms are concerned.

Certain changes similar to those occurring in digestion may take place in foods prior to ingestion In the ripening of certain fruits, such as the banana, starch is changed to devtrin and maltose through the action of amylase. Meats on storage undergo some self-digestion or autolysis. This process may be hastened for commercial purposes ("tenderizing") by the application of proteolytic enzyme preparations to meat cuts In the cooking of foods connective-tissue fibers are gelatinized and starch granules are broken up and some destrinization of starchy foods occurs. Cooking also increases the palatability of foods and in this way promotes the secretion of digestive juices.

Secretion of Saliva. The saliva is secreted by three pairs of glands—parotid, submavillary, and sublingual—reinforced by numerous small glands called huccal glands. These secretions vary in character. Thus in man the parotid saliva is watery and has a high digestive power. The secretions of the other glands are higher in mucin and more viscid. Ordinary saliva is a mixture of the secretions of these glands and shows considerable variations in composition in different individuals, and in the same individual at different times.

The secretion of saliva is governed by n set of nerve fibers cerebral in origin, together with fibers from the sympathetic nervous system. No hormone mechanism in salivary secretion is known. Ordinarily the secretion of saliva is the result of a reflex stimulation of the secretory nerves through a center in the medulla oblongata. Psychic stimuli, brought about by such influences as the thought of food, pass from the higher nerve centers to the secretory center and also give rise to secretion. The

results of Paylov on dogs show it to be rather difficult in certain cases to differentiate between the two types of stimuli Pavlov found that dropping several pebbles into a dog's mouth caused the flow of but one or two drops of saliva, but sand in the mouth induced a copious flow of a than watery saliva. Ice water caused no secretion, but acid or bitter solutions which the animal wished to reject caused a free flow of saliva. Dry food caused the secretion of a watery saliva; meat led to the flow of a more slimy secretion, such as would aid in the lubrication of this food for swallowing Drawing the attention of the animal to these foods, without actually giving them to bim, gave rise to similar secretions Thus, also, the pretense of throwing sand into the mouth of the dog gave rise to a profuse watery secretion.

The amount of saliva secreted by an adult in 24 bours has been vanously placed, as the result of experiment and observation, between 1000 and 1500 ml, the exact amount depending, among other conditions, upon the character of the food In the absence of obvious external stimuli, the rate of salivary secretion in the adult appears to he between 0.1 ml, and

09 ml per minute.

Composition of Saliva. Salivary composition depends on many factors: stimulation, diet, age, time of day, disease, etc To insure reproducihle, representative samples for analysis, the conditions for collection should include (a) a definite physiological, postabsorptive state (before breakfast), (b) no brushing of the teeth, rinsing of the mouth, or smoking prior to collection, (c) about a two-hour interval between arising and collection, including a 15-minute rest period immediately preceding collection Stimulation yields a relatively more dilute saliva, but the difference diminishes as collection is prolonged

Ordinarily saliva varies from weakly alkaline to weakly acid, the pH ranging approximately from 60 to 79 with an optimum pH of 66 There is evidence that normal individuals under 20 and without dental defects secrete a saliva with a pH between 70 and 72 No absolute correlation has been shown between salivary acidity and caries or other dental disorders However, lower pH values occur more frequently among earlessusceptible individuals. There is evidence that increased acidity of saliva is a late rather than early manifestation of the earies syndrome as compared with changes in the level of other salivary constitueats. Dental ero-ion is usually accompanied by greatly increased total salivary acidity. The acid secretion from the gingival crevice and the marginal gingival tends to dissolve enamel in the regions already predisposed as a result of unfavorable metabolic conditions

Saliva is a dilute secretion having an average specific gravity of 1.007 with about 0.7 per cent of solid matter, about 0.5 per cent being organic and 0.2 per cent morganic Of the organic matter about 0.4 per cent is protein, chiefly mucin, with small amounts of albumin and globulin. Other organic constituents are enzymes (chiefly salivary amylase), urea, uric and, chole-terol, vitamins, and phospholipide Average values have been reported of total nonprotein nitrogen 13 mg, urea plus ammonia natrogen 11 mg, and une acid 15 mg per 100 ml. These amounts ateraged 37 per cent, 76 per cent, and 40 per cent, respectively, of the corresponding constituents in blood Analyses of the saliva for these constituents possess a certain elimical value in the study of nephritis but are little used for this purpose. Normal saliva contains no glucose. Increases in cholesterol and in lipide phosphorus have been noted in certain dental disorders.

The normal composition of saliva is shown in the accompanying table

COMPOSITION OF NORMAL SALIVA*

Constituent	Λ ormal Range	
Acidity (pH)	60 - 79	
Titratable alkalinity (as 0 02 N HCl)	90 0 - 190 0 ml per 100 ml	
Ammonia N	2 0 - 10 0 mg per 100 ml	
Calcium, total, as Ca	40 - 80 mg per 100 ml	
Inorganic phosphate as P	10 0 - 25 0 mg per 100 ml	
Chloride as Cl	30 0 - 60 0 mg per 100 ml	
Carbonate as CO ₂	20 0 - 45 0 ml per 100 ml	
Protein	200 0 - 400 0 mg per 100 ml	
Cholesterol	2 5 - 9 0 mg per 100 ml	
Lipide P	0 05 - 0 20 mg per 100 ml	

*The values here given are based on saliva collected in a postabsorptive state and (with the exception of those for cholesterol and hipsde P) by stimulation Possible relationship to caries may be found in the following constituents ammonia calcium and phosphise (especially the adorbable fractions) carbonate cholesterol and hipsde P. Acknowledgment is made to Dr. Frances Krasnow for the data from which this table was compiled

Potassium thiocyanate ISSCN, is also generally present in the saliva to the extent of several milligrams per milliter. The significance of thiocyanite in the saliva is not known, it may possibly come from the ingested eyanides present in certain fruits and in tobacco smoke and from the breaking down of protein material Apoerythein a protein fraction which protects vitamin B₁₂ from digestive destruction is also present in saliva. This may be identical with the intrinsic factor of Castle.

Dam and his associates' have shown the following average content of vitamins in saliva (gamma (7) per ml) Thamine 0 007, nboffavin 0 05, macin 0 03, pyindovine 0 6, pantothenic acid (as Ca sali) 0 03 folic acid 0 0001, biotin 0 0008, ascorbic acid 2 4 vitamin K (as menadione) 0 015

The so-called tartar formation on the teeth is composed almost entirely of calcium and magnesium phosphates with some calcium carbonate minen epithelial cells and organic debris derived from the food. The calcium salts are held in solution as acid salts, and are probably precipitated by alkalimity caused by ammonia formation through bacterial action or through loss of CO. from the saliva. The various organic substances just mentioned are carried down in the precipitation of the calcium salts. There is evidence of increased subviry calcium in individuals suffering from excessive tartar deposition.

So-called salwary calcult may form in the gland or in the duct Their composition is similar to that of tartar This condition is known as stablishness

Salivary Amylase. The principal enzyme of the saliva is known as salivary amylase. The name pivalin, formerly used for this enzyme, is

Dam et al Intern Z l stammforsch 20 234 (1948)

now obsolete. Salivary amylase is an amylolytic enzyme. The enzyme catalyzes the hydrolytic splitting of starch, glycogen, and the dextrins into simpler molecules, the process being a progressive one with the disaccharide maltose as the ultimate end product.

The opinion has been confirmed that the salivary amylase activity varies with the composition of the diet and is highest on a diet which is

predominantly carbohydrate.2

The first product of the action of the amylase on starch is soluble starch, whose formation is indicated by the disappearance of the opales cence of the starch solution. Soluble starch gives a blue color with iodine; its formation from starch is not associated with the production of free reducing sugar (maltose), Further action of nmylase on starch apparently consists largely in the hydrolytic splitting of the second glycoside linkage from a free end of the long chain (straight or branched) of glucose residues which make up the starch molecule. This action produces the disaccharide maltose and a series of smaller polysaccharide molecules which are relatively ill-defined and which are known as the dextring.

The various dextrins differ in molecular size and complexity, depending upon the extent of amylase action. The higher members of the series resemble starch in giving a blue or purple color with iodine; as the molecule becomes smaller by the splitting off of maltose, a red color is given with iodine (erythrodextrins), and the lower members of the series give no color at all with iodine (achroodextrine). Thus during the action of salivary a-amylase on starch, free maltose is produced almost immediately and progressively increases in amount; at the same time the color reaction with iodine changes from blue through red to colorless. Both the increase in reducing sugar and the change in the rodine color reaction are used in following the course of action of amylase on starch. The changes described above may be represented as follows.

Starch	
Soluble starch	
Higher dextrins	Maltose
Erythrodextrins	Maltose
Achroodextrins	Maltose
Maltose	

These steps in the breakdown of straight and hranched glucose chains in amylose and amylopectin are illustrated in Figs 83 and 84 respectively

It must be borne in mind, however, that our knowledge of the course of starch hydrolysis by amylase is incomplete. This is due to the complicated nature of both starch (see Chapter 2) and amylase Most amylases appear to be mixtures of two enzymes, α- and β-amylase, which are said to

² Squires J Physiol., 119, 153 (1953).

produce respectively α - and β maltoses on starch by drolysis. According to one study on malt amylase, starch is completely by drolyzed to maltose by α - $+\beta$ amylase + a complement present with these in malt. Without the complement, 80 per cent of the starch may be split, leaving 20 per cent of residual dextrin with slight reducing power α -Amylase alone leaves about 60 per cent of dextrins giving no color with iodine, while β amylase alone leaves about 60 per cent of dextrins giving a blue color with iodine. The amylase alone leaves about 40 per cent of dextrins giving a blue color with iodine in malt extract. Malt extract kept for several days at 0° C and at pH 3 6 contains only β amylase α Amylase is obtained by extraction of ungerminated harley with water

Salivary amy lase acts in alkaline, neutral, or faintly acid solutions. The optimum acidity is pH 6.6. Amy lase is destroyed at acidities greater than



FIO 83 ACTION OF α-AMYLASE ON AMYLOSE (O)
GLUCOSE UNIT (O-O-) α 1 4-GLUCOSIDIE LINKAGE
(1) α-AMYLASE ACTION
P Betrifeld Adv Entymol 12 379 (1951)

pH 4 or at concentrations of free HCl greater than 0 0005 per cent. By sufficiently increasing the alkalimity of the salvar the action of the salvary amylase is inhibited. Salvary amylase has been crystallized.

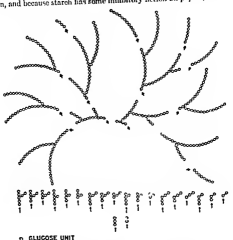
Electrolytes have an important influence upon the action of amylases For example, Rockwood has shown that Cl Br, and NO₂ ions have a pronounced stimulating action upon sulvary amylase Removal of chlorides from saliva by dualysis renders the amylase inactive Amino acids particularly asparagine, have an accelerating action Salts of the beavy metals such as silver and mercury inhibit because they combine with the citizene, which is apparently of a protein nature

Because of its sensitivity to acid, salivary amylase ceases to act in the stomach as soon as the gastric contents show throughout the presence of free hydrochloric acid. Since, however, the amylolytic activity of human saliva is very great, an appreciable digestion of starch may occur during the period of mastication and swallowing. Furthermore, the food which is swallowed is not immediately mixed in its entirety with gastric juice and the protein of the food has a certain binding power for free acid, so that a certain interval may intervene between the entrance of the food into the stomach and the destruction of the amylase. This period varies much in different animals and depends also on the size and charac-

¹ Meyer et al Helv chim. Acta 31 2158 (1949)

ter of the meal. In cert un experiments on normal men Bergeim found that salivary digestion might continue for 15 to 30 inmutes, and that in meals with bread and mashed potitoes there was a conversion of the starch to maltose of about 60 and 75 per cent respectively

The digestion of the starch of foods decreases their bulk, and for this reason, and because starch has some inhibitory action on pepsin, salivary



REDUCING END GROUP

6-GLUGDSIDIG LINKAGE

g-1.4-GLUGD SIDIG LINKAGE

Fig 84 Action of a Antlabe on Antlopectin

Upper portion Formation of sodine reacting dextrins of medium molecular weight I ower portion Possible structures of limit dextrins (heptasaccharides, hexasac charides etc.)
P Bernfeld Ads Enzymol 12 379 (1951)

digestion may have some favorable effect on protein digestion in the stomach The action of salivary amylase is not, however, essential, since the pancreatic juice contains a powerful amylase

The saliva of rodents is amylolytic Hog saliva contains amy lase, but much less than human saliva. The saliva of carnivorous animals is free from amylase, as is also that of herbivorous animals

Maltase is found in traces in saliva Maltase splits maltose into glucose but the amount of such digestion in the mouth is very slight. It is claimed that dipeptide- and tripeptide-splitting enzymes are present in saliva. This action in at least some eases is due to bacteria and in any case is not of directive importance.

Mucin. Mucin gives saliva its viscosity It is a glycoprotein, insoluble in water or dilute acid hut soluble in dilute alkali. When precipitated, as upon the teeth, mucin forms with alkali a slippery mass which dissolves but slowly. It is therefore removed from the teeth with difficulty and may furnish a nucleus for the deposit of other substances. Mucin may he precipitated from saliva by dilute acid or by alcohol. It is an acid substance existing in saliva as the potassium salt. It gives the usual protein color reactions, but is not coagulated by heat in neutral solutions. It is precipitated by saturating with ammonium sulfate. On hydrolysis it yields besides protein a mucoitin sulfurie acid which on further decomposition gives sulfurie acid, acetie acid, glicuronic acid, and glucosamine.

Microscopical examination of saliva reveals epithelial cells, salivary corpuscles (white blood cells?), mucus food debris, and numerous microorganisms from the true bacteria, higher bacteria, fungi, and protozoa groups Pus cells and red blood cells may be evident in pathological conditions of the mouth

EXPERIMENTS ON SALIVA

A satisfactory method of obtaining the saliva necessary for the experiments which follow is to chew a small piece of pure paraffin way, thus stimulating the flow of the secretion, which may be collected in a small beaker. It must be remembered in this connection that paraffin stimulated saliva is quite different in some respects (e.g., p.H.) from ordinary saliva. Filtered saliva should be

used in every experiment except the microscopical examination and the quantitative determination of amplase activity

I Microscopical Examination Examine a drop of unfiltered saliva microscopically, after staining with methylene blue, and compare with Fig. 85



Fig 85 Microscopic Constituents of

2 Reaction Test the reaction to iltmus or other suitable indicator paper Estimate the approximate pH a Epithelial cells, b salivary corpuscles, c fat drops d leukocytes, c, f, g bacteria, h i k fission fungi

of the saliva Measure 2 mi of fresh saliva into a small test tube Add 10 drops of an indicator solution suitable for measuring pil at the estimated pil of the saliva 4 Compare with 2-mi portions of pil standard solutions4 treated with the same a mount of indicator in similar tubes. The standards should differ by

⁴ See Chapter 1 for the preparation and use of indicators standards and the composition

0.2 pH unit. Determine the pH of the saliva. The comparison is best made in a comparator block, with a tube containing saliva only placed behind the standard, and a tube of plain water behind the saliva-indicator tube.

- 3. Test for Mucin. To a small amount of sallva in a test tube add 1 to 2 drops of dilute acetic acid. Mucin is precipitated.
- 4. Biuret Test. Render a little saliva alkaline with an equal volume of NaOli and add a few drops of a very dilute (2 to 5 drops in a test tube of water) copper sulfate solution. The formation of a purplish-violet color is due to mucin. This reaction is given by protein material and simply indicates that mucin is a protein.
- 5. Preparation of Mucin, Pour 25 ml. of saliva Into 100 ml. of 95 per cent alcobol, stirring constantly. Cover the vessel and allow the precipitate to stand at least 12 hours. Pour off the supernatant Ilquid, collect the precipitate on a filter, and wash it, in turn, with alcohol and ether. Finally dry the precipitate, remove it from the paper, and make the following tests on the mucin: (a) Test its solubility in water, dliute acid, and dilute alkali; (b) Millon's reaction; (c) dissolve a small amount in NaOII, and try the bluret test on the solution; (d) boil the remainder, with 10 to 25 ml, of water to which 5 ml. of dilute IICl has been added, until the solution becomes brownish. Cool, render alkaline with soild sodium carbonate, and test by Benedict's solution. Reduction should take place. Why? Does the unhydrolyzed mucin reduce Benedict's solution?

Mucin may also be prepared from salivary glands.

- 6. Viscosity Test. Place filter papers in two funnels, and to each add an equal quantity of starch paste (5 ml.). Add a few drops of saliva to one lot of paste and an equivalent amount of water to the other. Note the progress of filtration in each case. Why does one solution filter more rapidly than the other?
 - 7. Test for Nitrites. Add 1 to 2 drops of dilute li: SO, to a little saliva and stir thoroughly. Now add a few drops of a freshly prepared potassium lodide solution and some starch paste, Nitrous acid is formed which liberates lodine, causing the formation of a blue color with the starch.
 - 8. Thiocyanate Test: Solera's Reaction. This test depends upon the liberation of lodine through the action of this cyanate upon lodic acid, Molaten a strip of starch paste-iodic acid test papers with a little saliva. If thiocyanate is present the test paper will assume a blue color, owing to the liberation of iodine and the subsequent formation of the so-called lodide of starch.
 - 9. Digestion of Starch Paste. To 25 ml. of starch paste in a small beaker, add 5 drops of saliva and stir thoroughly. At intervals of a minute remove a drop of the solution to one of the depressions in a test tablet and test by the lodine test. At the same time add 3 drops of the mixture to one of a series of test tubes set up with 5-ml. portions of Benedict's reagent. The opalescence of the starch solution should soon disappear, indicating the formation of

See Appendiz,

To obtain a closer approximation of the reaction of saliva in the mouth, draw the samrie into a riget directly from the mouth (without previous chewing of paraffin), and dilute with the indicator in the test tube under a thick layer of sameral oil (Krasnow, Oblatt, and hajlan J Dental Research, 15, 367 (1936).)

soluble starch which gives a blue color with lodine. The soluble starch should soon be transformed into erythrodestrin which gives a red color with lodine, and this in turn should pass into achroodestrin which gives no color with lodine. This is called the achromic point. When this point is reached, complete the Benedict tests by placing all the tuhes in a boiling water bath for 3 minutes, and note the degree of reduction in each tube. Tabulate your results with the lodine and Benedict tests in parallel columns. Also perform a phenylhydrazine test for maltose (the osazone crystals may not have the typical appearance of maltosazone). A positive Benedict test may be obtained while the solution still reacts red with lodine, inasmuch as some maltose is formed from the soluble starch coincidently with the formation of the erythrodestrin. How long did it take for a complete transformation of the starch? Saliva from different individuals may vary markedly in amylolytic power. For a graphic representation of the above changes see p. 352.

- 10. Separation of the Products of Salisary Digertion. To 25 mi., of 1 per cent starch paste in a small beaker add I mi. of saliva and stir thoroughly. At intervals of one minute test a drop of the mixture by the iodine test. If the hine color persists after five minutes add another I mi. of saliva. When the mixture reacts red with lodine, indicating that erythrodextrin has been formed, add 100 mi. of 95 per cent alcohol. Allow to stand until the white precipitate has settled. Filter, evaporate the filtrate to dryness on a water bath, dissoive the residue in 5 to 10 mi. of water, and try Benedict's test and the phenylbydrazine reaction. On the dextrin precipitate try the lodine test.
- 11. Influence of Temperature. Into each of four tubes place about 5 ml. of 0.2 per cent starch paste. Immerse one tube in a beaker containing crushed ice, keep a second at room temperature, and place a third in the incubator or the water bath at 40° C. (If the temperature of the hath or incubator is allowed to rise to 70° C. or over the enzyme is destroyed and no digestion takes place.) Now add to the contents of each of these three tubes 5 drops of saliva and shake well; to the contents of the fourth tube add 5 drops of boiled saliva. Test frequently by the lodine test, using the test tablet, and note in which tube the most rapid digestion occurs. Explain the results.
- 12. Estimation of Amylare in Salina. Pipet exactly 1 ml. of unfiltered saliva Into a 100-ml. cylinder. Dilute to the 100-ml. mark, and mix well. Pipet 5 ml. of 1 per cent soluble starch Into a test tube. Add 2 ml. of 1 per cent NaCi solution and 2 ml. of a buffer solution of pH 6.6° and put In a water bath maintained at 38° C. Prepare a series of 10 test tubes, each containing 2 ml. of a light yellow lodine solution. Now add 1 ml. of diluted saliva to the starch mixture and return to the bath at once. Record the time of this addition. At the end of each minute of digestion, with a pipet remove 2 drops of the mixture and add to one of the tubes of lodine solution. Record the time when no change in color appears in the lodine solution (achromic point). If this time is less than 5 or more than 20 minutes, repeat, using a different dilution of saliva such as will give a digestion time of about 10 minutes. Thus If 3 minutes are required, dilute 30 ml. of the original diluted saliva to 100 ml., and use 1 ml. of this for the test.

One unit of amylase may be considered to be the amount required to digest 5 mi. of 1 per cent soluble starch to the achromic point in 10 minutes under the conditions of the test. The number of units of amylase in 1 mi. of the sailva tested will equal 100 (or whatever the dilution of the original sailva might be) × 10 + the number of minutes to the achromic point. From 100 to 150 units are frequently found.

- 13. Influence of NaCl on Salmary Amylase. Repeat the preceding experiment hut replace the NaCl solution by water. A longer time should be required, showing that NaCl accelerates the action of salivary amylase. If saliva be dialyzed free from chlorides it hecomes inactive.
- 14 Influence of pH on Salisary Amylase. In Exp. 12 the solution had a pH of 66. Replace the phosphate buffer of this experiment hy another huffer of pli 5.8 and in a second experiment use a buffer of pli 8.0.7 Which gives digestion in the shortest time and is hence nearest the optimum pli?
- Excretion of Potassium Iodide. With the aid of a glass of water, ingest a small dose of potassium iodide (0.2 g.) contained in a gelatin capsule, quickly rinse out the mouth with water, and then test the saliva at once for lodine. This test should be negative. Make additional tests for iodine at twominute intervals. The test for sodine is made as follows: Take 1 ml. of dilute NaNO: solution and 1 ml. of dilute Ii, SO, In a test tuhe; add a little sallva directly from the mouth, and a small amount of starch paste. The formatioo of a blue color signifies that the potassium lodide is helng excreted through the salvary glands. Note the length of time elapsing between the ingestion of the potassium lodide and the appearance of the first traces of the substance in the saliva. If convenient, the urine may also be tested at 15- or 30minute intervals after ingestion of the iodide. The chemical reactions taking place in this experiment are indicated in the following equations:
 - (a) 2\sNO₂ + II₂SO₄ → 21INO₂ + Na₂SO₄
 - 2K1 + H,SO, -> 2HI + K,SO,
 - (c) $2HNO_{2} + 2HI \rightarrow I_{1} + 2H_{2}O + 2NO_{2}$

Inasmuch as iodide is absorbed from the stomach very slowly, If at all, but is very rapidly absorbed when it enters the intestine, the rapidity of appearance of the lodide in the saliva is an index of the rapidity with which the drug leaves the stomach, which depends on the motor activity of the stomach, the amount of food therein, etc. By drinking a glass of water with the iodide a more rapid result is obtained,

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⁷ See Chapter 1.

⁵ thatead of this muxture a few drops of ItNO: possessing a vellowish or brownish color (containing ItNO:) may be employed

14

Gastric Digestion

Following mastication the food is carried by peristaltic movements of the esophagus to the stomach. Here it undergoes further mechanical disintegration and chemical changes primarily in the protein constituents. The food thus treated is in a better condition to be handled by the intestines, to which it is passed on in small portions at a time, and in which digestion is completed.

Spallanzam (1783) found that gastric juice dissolved meat, and so gastric digestion He noted vise that the juice was acid, but the nature of this acid and of the active agent

pepsin was not demonstrated until later

A great advance in our knowledge of gastrie digestion, particularly in man, was made through the observations of Beaumont on his patient Alcus St Martin who in 1822, following a guisbot wound was left with an opening from the stomach through the abdominal wall to the exterior Through this fistula Beaumont found it possible to follow the course of gristric digestion with different foods and under varying conditions of health, and to obtain pure gastric juice for digestion experiments outside the body

Pavior extended our knowledge particularly through the development of an operation by means of which he created in dogs a small stomach pouch separate from the main stomach and opening to the exterior, so that the secretion in the small pouch could be studied without interfering with processes in the stomach proper. In this way the influence of different foods and of other factors on gastries exerction could be studied. The development of the small stomach tube which could be retained in place throughout the period of gustrie digestion and allowed aspiration of the stomach contents at any time, has given additional information of physiological and clinical value.

Secretion of Gastric Juice There is a slight continuous secretion of gastric juice into the empty stomach As a result there is almost always present in the stomach before meals about 50 ml of secretion which is called the residuum Gastric secretion is governed by many factors. At

least three phases are recognized

(1) PSYCHIC PHASE Following the presentation of food and before any food reaches the stomach, there is a psychic secretion of gastric juice. This is induced by the sight, taste, smell or thought of food. Try has called this the "cephalic phase."

(2) Gastric Phase Pollowing this, with the passage of food into the stomach, there occurs the gastric phase of gastric secretion due to the

local chemical action of such substances as protein digestion products or meat extractives and some mechanical action due to friction and distention of the stomach by food A substance, gastrin, has been isolated from gastric mucosa (Edkins) When injected into the blood this mark edly stimulates gastric secretion. It is believed to be a hormone of gastric secretion, i.e., a substance liberated from the gastric mucosa in the presence of food and passing by way of the blood to the acid secreting cells, stimulating them to action Histamine the decarboxylation product of the amino acid bistidine is also a powerful stimulant of gastric secretion in addition to its well known effect on blood pressure (capillary dilatation) The question whether or not gastrin is identical with hista mine was unanswered for some years. They are now believed to be separate substances which possess similar action 1 The chemical structure of histamine is shown on p 1037 The structure of gastrin is not known Clinical use is made of the secretagogue effect of histamine, alcohol, and caffeine

(3) INTESTINAL PHASE A third intestinal phase of gastrie secretion is brought about through the action of protein digestion products and other food substances in the intestines Whether these substances act by liberat ing a hormone from the mucosa, or are absorbed and themselves act upon the gastric cells is not yet clear Undigested fat in the intestine inhibits gastric secretion, apparently by liberating from the mucosa a hormone called enterogastrone which depresses the action of the gastric cells This chalone also inhibits gastric movements and was for a time considered useful in the treatment of gastric ulcer However evidence of its in effectiveness has accumulated For example Wollum and Pollards found no significant alteration in the secretory or motor patterns from its use and Bone' has reported that it has little or no value Chinical research has indicated that the chemical substances called Banthine Pro-banthine and Prantal are useful in the treatment of ulcer (see p 383) Urogastrone a substance similar in action to enterogastrone and found in the urine 18 apparently a metabolic derivative of enterogastrone

Water has a stimulating action on gastric secretion and the drinking of considerable water has been shown to improve the utilization of various foods Nor has the drinking of water with meals by normal individuals been shown to be undesirable 'The influence of different foods on gastric

secretion is discussed later

The study of gastric secretion and the gastric mucosa in nutritional deficiencies has received some attention. Thus in canine blacktongue (the closest animal counterpart to human pellagra) a mild pallor of the gastric mucosa was observed consistently upon gastroscopic examination This was accompanied by anemia loss of weight and decreased muscular tone of the stomach wall The tonus of the stomach returned to normal as early as six to seven days after the institution of therapy. There was no

Friedman and King Federation Proc. 6 107 (1947) Wollum and Pollard Gastro-nterology 17 535 (1951)

^{*} House and Controlled
change in acid gastric secretion during the disease or following macin therapy The therapeutic effect of thiamine was also negative Shapiro and his co-workers found atrophic gastritis to he somewhat more common in nutritional deficiencies, than in a control group of ten other patients However, no significant changes in the gastric mucosa were observed after treatment with large doses of thiamine, macin, riboflavin, pantothenic acid, p-aminohenzoie acid, and vitamin A In two cases the atrophic changes disappeared after choline chloride therapy

Composition of Gastric Juice. Normal gastric juice is a thin, hight-colored fluid which is acid in reaction and has a specific gravity averaging about 1 007 It contains about 0.5 per cent of solid matter which is made up principally of sodium chloride, potassium chloride, earthy phosphates, mucin, and the enzymes pepsin and gastric lipase The acidity of the gastric juice is due to free hydrochloric acid. The gastric nuce is a composite secretion from at least three different types of cells in the gastric mucosa, these are (1) the parietal cells, (2) the chief cells, and (3) the mucous cells There is good evidence that the parietal cells furnish the hydrochloric acid of gastric juice, the chief cells supply pensin and possibly other enzymes, and the mucous cells secrete mucin Babkin' claims that the secretory activity of the various types of gland cells should not be considered en masse but rather that "various nerves

or chemical agents stimulate or inhibit each set of secretory ele-

ments separately " This view is also accepted by others "

Apoeruthein, a protein fraction, also occurs in gastric juice 9 It has been suggested that this substance may he identical with the intrinsic factor of Castle, which protects vitamin B12 (the extrinsic factor) from directive destruction Permicious anemia is characterized by a deficiency of the intrinsic factor Vitamin Bio is relatively ineffective when given orally to such patients, but is highly effective when administered parenterally

It is helieved that the parietal cell secretion is essentially an isotonic solution consisting largely of hydrochloric acid (about 160 millieguivalents per liter) and potassium chloride (about 7 milliequivalents per liter) The acidity of the parietal-cell secretion corresponds therefore to a solution 0 16 N in hydrochloric acid, or containing 0 5 to 0 6 per ceot hydrochloric acid This maximal acidity, which is apparently constant and independent of the rate of secretion, is lowered somewhat as soon as the parietal-cell secretion becomes admixed with the slightly alkaline secretions from the chief cells and mucous cells. These latter secretions contain a high concentration of neutral chlorides as n result of which the acidity is reduced to a greater extent than is total chloride content. The acidity may also he lowered by regurgitation of alkaline fluid from the intestine and hy ingested food, so that the actual acidity of gastric juice as collected usually varies between 0 05 and 0 1 N (0 18 to 0 36 per cent hydrochloric acid) The acidity of the gastric juice is usually expressed in terms of the number of milliliters of 0 1 N sodium hydrovide required to

Shapiro et al., ibid 2 121 (1944)
 Babkin 4m J Digest Diseases 5 10" (1937) 8 46" (1938)
 Thomas J Am Med Assoc 120 "35 (1942)
 Ternberg and Eakin J 4m Chem Soc 71 3358 (1949)

neutralize 100 ml. of gastrie juice; this is obviously equivalent to the number of millithers of 0.1 N hydrochloric acid present in 100 ml. of gastrie juice, which corresponds numerically with the concentration of and expressed in terms of milliequivalents per liter. In clinical practice this value is sometimes called the degree of aridity; thus a gastrie juice containing 60 milliequivalents of acid per liter, or requiring 60 ml. of 0.1 N alkalt to neutralize 100 ml., is said to have an acidity of 60 degrees

The hydrochloric acid of the gastric juice forms a medium in which the pepsin can most sati-factorily digest the protein food, and at the same time it acts to some extent as an anti-eptic or germicide which prevents putrefactive processes in the stomach. When the hydrochloric acid of the gastric juice is diminished in quantity (hypoacidity) or absent, as it may be in many cases of functional or organic disease, there is no check to the growth of microorganisms in the stomach. There are, however, certain of the more resistant spores which even the normal acidity of the gastric juice will not destroy. A condition of hypoacidity may also give rise to fermentation with the formation of comparatively large amounts of such substance as lactic acid and butyris acid

When free hydrochloric acid comes in contact with protein, as in the food, a reaction occurs with the formation of protein hydrochloride. This was formetly called "combined hydrochlorine cacid," but this term is so indefinite in its connotation that it should be abandoned (see p. 380). The formation of protein hydrochloride considerably raises the pH of gastric contents, since protein hydrochloride as much less highly ionized acid than is hydrochloric acid itself. The reduction in hydrogen-ion concentration resulting from the formation of protein hydrochloride may permit processes which are acid-sensitive to proceed during gastric digetion, such as bacterial action or the action of salivary amylase (see p. 353)

Origin of Castric Acid. The mechanism whereby the stomach produces a secretion which is about three million times more acid than the blood is not known Many attempts have been made to solve this problem, and various theories have been proposed, none of which has received universal acceptance

The currently favored view for the formation of acid by the parietal cells is based upon the discovery by Davenportion that the enzyme carbonic anhydrase (see Chapter 21) is present in large amounts in the parietal cells, and is relatively absent from the other cells of the gastric nuccoa. This enzyme also occurs in red blood cells. It has been crystallized and found to contain 0.2 per cent zine 11 Potent inhibitors of carbonic anhydrase (like certain sulfa drugs) also depress hydrochloric acid secretion in the living animal 12 Carbonic anhydrase entalyzes the hydration of carbon dioxide to carbonic and, which dissociates in solution to yield hydrogen ions and brearbonate ions.

 $CO_2 + H_2O = H_2CO_3 = H^+ + HCO^-$

²⁸ See Davenport Gastroenbrology 1, 283 (1943), Gray Dol. 1, 390 (1943)
²⁸ Summer and Somers Chemistry and Methods of Enzymes 3rd ed. New York, Academic Press Inc., 1976.

¹¹ Janowitz Colcher, and Hollander Trans NY Acad Sci., 15, 54 (1952)

If this reaction is pictured as occurring within the parietal cell, the carhon dioxide coming from metaholic processes, the hydrogen ions may be visualized as heing secreted, along with an equivalent number of chloride ions, into the stomach while the hierarhonate ions enter the hlood

Here hicarhonate ion replaces chloride ion, which diffuses into the parietal cell and is available for excretion along with hydrogen ions into the gastric juice. Thus according to this theory, the hydrogen ions of the gastric juice come from carhonic reid, and the chloride ions are derived from the blood. The replacement of blood chloride by hicarhonate should raise the pH of the blood, and in fact it has been shown that blood leaving the stomach during active gastric secretion is significantly more alkaline than the entering blood thus accounting for the "alkaline tide" (see p. 7841).

It is an interesting consequence of this theory that the extra bicarbonate of gastric venous plasma as compared to gastric arterial plasma is the result of the direct entrance of bicarbonate as such into the plasma and not the result of the entrance of anhydrous carbon dioxide into the blood followed by its hydration to carbonic acid in the red cell and diffusion from the red cell into the plasma, processes which are known to account for the extra hicarbonate of venous plasma over arterial plasma in other parts of the hody Now in ordinary venous plasma the increase in plasma bicarbonate due to diffusion of bicarbonate from the red cell into the plasma is associated with a chloride shift (see Chapter 24), chloride ions leaving the plasma and entering the red cell In gastric venous plasma the increased plasma hicarhonate hrings about a reversed chloride shift by diffusion into the red cell at the expense of chloride ions which diffuse out into the plasma. This has been cited as further evidence concerning the role of carbonic anhydrase in the formation of gastric acid It should be remembered, however, that though this theory accounts for the production of acid by the parietal cell, no direct evidence concerning the actual mechanisms of formation and secretion is as yet available, and other theories12 are not as vet untenable

Pepsin. The most characteristic of the enzymes of the gastric juice is the proteolytic enzyme pepsin. Pepsin is a representative of a large group of enzymes, many of which are found in the gastrontestinal tract and all of which catalyze the hydrolytic spbtting of the peptide bond,—CO—NH—to produce a free amino and a free carboxyl group. Within this group of peptide-splitting enzymes, two general by pes may be distinguished (Bergmann). (1) the protemases or endopeptidases, and (2) the peptidases or exopeptidases. The endopeptidases not upon peptide linkages in both the central portion and the terminal portion of a polypeptide chain, the exopeptidases spht peptide linkages in the terminal portion of the chain only. Differences between the various endo- and exopeptidases of the gastronitestinal tract are attributed largely to differences in the type and location of the amino acids united in the peptide bond, as will be evident in this and sub-equent clumpters. According to this classification, pepsin is an endopeptidase, since it can act upon peptide linkages.

[&]quot; See Hollander Gastroenterologs 1 401 (1943) Fe leration Proc 11 "06 (1952)

within the large protein or polypeptide molecule as well as upon synthetic

peptides (see below)

Pepsin is apparently formed by the action of the hydrogen ions of the gastric juice on a precursor or zymogen, called pepsinogen, originating in the chief cells of the gastric mucosa Both pepsin and pepsinogen are proteins and have been prepared in erystalline form,14 crystalline pepsin appears, however, to be a mixture of enzymes rather than a single substance Pepsinogen is more resistant to alkalı tban is pepsin. It does not elot milk at pH 5 nor liquefy gelatin at pH 47, but pepsin is active under these conditions The formation of pepsin at pH 46 appears to be an autocatalytic reaction, i e , the pepsin as it is formed acts upon further pepsinogen to yield still more pepsin. Since pepsin is known to aet only on peptide linkages with a specific amino acid configuration (see below), it would appear that the activation of pepsinogen may involve the breaking of such linkages. The isoelectric point of pepsin is at such a low pH that it has not been accurately established

Pepsin contains phosphorus It is denatured and coagulated by heat, with the loss of peptic activity running parallel with the degree of denaturation It is also inactivated and denatured in alkaline solutions (pH 10) In such cases there is some return of activity on acidification and standing, thus there has been a reversal of the denaturation of the pepsin protein Pepsin is digested by trypsin in solutions more alkaline than pH 55 In more acid solutions trypsin is digested by pepsin It has been shown that when pepsin is mixed with egg albumin in the pH range 3 1 to 4 2 a precipitato is formed which is not denatured

Pepsin acts very well at 40° C The optimum pII is about 20, but this has been shown to be more related to the ionization of the substrate protein than to any effect on the enzyme itself A variety of mineral and organic acids may be used to attain this pH with practically identical effects on the enzyme activity At pH 4 its activity is very slight and at pH 5 it is stable but mactive At pH values of 6 0 or greater it is unstable, and above pH 8 it is rapidly destroyed by OH ions

Products of Peptic Hydrolysis. The gastrie acid acting on food proteins at body temperature produces denatured proteins ("acid metaprotein") Pepsin acts upon such denatured protein as well as upon native protein with the formation largely of protein derivatives of relatively low molecular weight (not over 1,000, according to Tischus and Eriksson-Quensel16), such split products belong to the ill-defined class of the proteoses and peptones In the normal time of gastrie digestion, substances of this nature appear to be the chief end products of peptie action Free tyrosine is also a frequent product of peptic digestion. The nature of the action of pepsin has been considerably elarified by the work of Bergmann, Fruton, and associates They have studied the action of crystalline pepsin

¹⁴ For discussion of crystalline pepsin see pp 305 and 330 For preparation of crystalline pepulogen see Herriott and Northrop Science 83 469 (1936) and Herriott J Gen Phymol. 21 501 (1938)

¹⁴ lasnoff and Bull J Biol Chem 200 619 (1953) " Buchem J 33, 1752 (1939)

on synthetic substrate peptides of known chemical constitution. All synthetic substrates which were hydrolyzable by pepsin contained either tyrosine or phenylalanine in the molecule, with the peptide linkage specifically involving the amino group of these amino acids. If the tyrosine or phonylalanine was at the end of the peptide chain, hydrolysis proceeded more rapidly and the free amino acid was liberated, but peptic action was not limited to the presence of the amino acid at the end of a chain. Thus one may conclude that pepsin acts on peptide linkages associated with the presence of tyrosine or phenylalanine (although the possibility of other amino acids being concerned in peptic action is not necessarily excluded); the end product of the action will depend upon the relative nosition of these amino acids in the long chain of amino acid residues in the protein substrate. Pepsin acts upon practically all native proteins with the exception of keratin and the protamines; its lack of action in the case of keratins is believed to be due to the close packing of the polypentide chains in the keratin molecule; in the case of protamines, to their deficiency in tyrosine and phenylalanine.

In testing peptic activity at pH from 1 to 2.85 Currie and Bull' report that pensin acts as a proteinase at the lower pH and splits fewer pentide linkages per mole of substrate thus yielding fragments of relatively high molecular weight. On the other hand, at a higher pH it acts as a peptidase and yields fragments of smaller molecular weight. As to the influence of temperature it was found that at 30° C. or helow the digestion products possessed relatively low molecular weights whereas at temperatures hetween 35° C, and 50° C. the products of hydrolysis were of high molecular weight. The conclusion can therefore be drawn from these experiments that pepsin acts very efficiently at pH 2.85 and at about 30° C.

Rennin. Rennin is what is known as a milk-curdling or protein-coagulating enzyme. It is a proteinase which, acting upon the cascin of milk, is believed to form first a soluble paraeasein and a peptone-like body. In the presence of ionized calcium salts there is then formed an insoluble calcium paracaseinate which separates out as a curd. Rennin is commonly obtained from the mucosa of the fourth stomach of the calf and is used to curdle milk in cheese-making.

The matter was the subject of a long controversy, but it appears now to be clear that the enzyme, remain, is an entity quite distinct from pepsin. It has very high milk-curdling power but practically no protein-digesting activity. In certain animals the curdling of milk is caused by pepsin, Pcpsin can coagulate milk in practically neutral solution as contrasted with the high acidity required for its action on proteins in general.

Rennin acts best at a pH of 6.0 to 6.5 and at a temperature ci about 45° C. Tauber and Kleiner have obtained a preparation curdling 4,550,000 times its weight of milk at pH 6.2 in 10 minutes at 40° C. The preparation has an isoelectric point of 5.4. It is apparently a diffusible proteose containing sulfur and is not congulated by heat. It does not give the Millon and Hopkins-Cole tests. Rennin appears to exist in the calf's mucosa as prorennin, which is activated by the gastrie acidity.

¹⁷ Currie and Bull: J. Biol. Chem., 193, 29 (1951).

to the evacuation time and the highest total acidity after the ingestion of certain common foods by normal men. In the tests here summanized 100 g portions of food (unless otherwise stated) were fed to normal men and the grattic reponse determined by the fractional method (see p. 383). It will be noted in general that foods such as meats which are high in

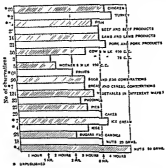


FIG 90 EVACUATION TIMES OF COMMON FOODS Hawk Pehlum and Bergeim Am J Med Set. 171 359 (1926)

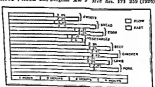


FIG 91 THE PLACUATION TIMES OF FAST AND SLOW STOMACHS

Hawk, Reblum and Bergeim Am. J Med Ses. 171, 359 (1928)

protein, and for which gastrie digestion is hence of the greatest importance, remain longest in the stomach (3 to 4 hours) and give rise to the highest acidities (120 or higher) Foods low in protein, such as fruits and many vegetables leave the stomach soon (1 5 to 2 hours) and give rise to much less secretion Foods intermediate in protein content, such as creal foods, show intermediate acidities and emptying times. Wilk has considerable buffer action Water leaves the stomach rapidly. The relationship of the average evacuation times of the various foods.

is shown graphically in Fig. 90, whereas the variation in the evacuation time of the same food by fast and slow stomachs is shown in Fig. 91.

COLLECTION OF HUMAN GASTRIC JUICE

Have one or more volunteers from the class take the Rehfuss stomach tube as directed on p. 384. The subjects must omit breakfast if the tube is taken in the morning, or luncheon if taken in the afternoon. Empty the stomach (see

pp. 384 385) and, with the tube still in place, allow each subject to drink 250 ml. of water. The water will stimulate the flow of gastric juice and will itself quickly leave the stomach. In many instances fairly concentrated gastric juice may be obtained from the stomach in from 30 to 45 minutes after the introduction of the water, Remove this gastric juice according to procedure outlined on p. 386. For the composition of human eastric juice see p. 377. See also Exp. 9, p. 373. If thought desirable, the eastric juice resulting from psychical stimulation (see p. 373) or following histamine injection (see p. 385) may be collected instead of that following the chemical stimulation of water. (Curves showing the stimulatory power of water are given in Fig. 92.)

PREPARATION OF ARTIFICIAL GASTRIC JUICE

1. From Pig's Stomach. Dissect the mucous membrane of a pig's stomach from the muscular portion and discard the latter. Divide the mucous mem-

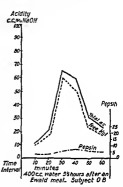


FIG 92 CURVES SHOWING STIMU-LATORY POWER OF WATER Betgerm Rebluss and Hawk J Biol Chem 19, 345 (1914)

brane Into two parts (four-fifths and one-fifth). Cut up the larger portion, place it in a large-sized beaker with at least 4 volumes of 0.4 per cent hydro-chloric acid, and kep at 38° to 48° C. for at least 24 hours. Add more HCl as needed to keep the mixture acid to Congo red paper; otherwise putrefaction may occur. Filter off the residue, consisting of nuclein and other substances, and use the filtrate as an artificial gastric juice. This filtrate contains pepsin and the products of the digestion of the stomach tissues; i.e., denatured protein, proteoses, and pentones.

2. From Commercial Pepsin. Dissolve 750 mg, of U.S.P. pepsin in 100 ml. of 0.1 N hydrochloric acid.

PREPARATION OF A GLYCEROL EXTRACT OF PIG'S STOMACH

Take the one-fifth portion of the mucous membrane of the pig's stomach not used in the preparation of the artificial gastric juice, cut it up finely, place it in a small-sized beaker, and cover the membrane with glycerol. Stir frequently and allow to stand at room temperature for at least 24 hours. The glycerol will extract the pepsinogen. With a pipet or by other means, sepa-

The Action of Gastric Juice on Milk A comparison between the large tough curds formed in the stomach from cow's milk and the small soft curds formed from human milk is shown in Figures 86 to 89, below is

Gastric Lipase A third enzyme present in gastric juice is a fat split ting enzyme. It possesses but shight activity when the gastric juice is of



Fig. 86 Curd of Human Milk Five Minutes after Indestion of Milk. Beginning of curd formation Onehalf actual size

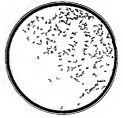


Fig 87 Curd or Human Mile Ten Minutes after Indestion or Mile. Maximum curl formation Onehalf actual size



F1: 88 CURD OF COWS MILK REGUE CITATED 10 MINUTES AFTER INGES-TION OF 500 MIL OF WHOLE MILK One-half actual size



Fig 89 Curd of Cow 8 Mile Regur-GITATED 25 MINUTES AFTER INGES-TION OF 500 ML OF WHOLE MILE One half actual size

normal acidity, but evinces its action principally at such times as a gastric juice of low acidity is secreted either from physiological or pathological cause It thus may be of importance in the young animal where gastric acidity is considerably lower than in the adult. The digestion of fat in the stomach is however at most of but slight importance as compared with the digestion of fat in the intestine through the action of the

ss Bergeim Fryard Relfuss and Hawk Am J I has of 48 411 (1919)

lipase of the pancreatic juice (see p 397). The presence of lipase in the gastric lumen, like that of trypsin, is probably due to regurgitation of intestinal contents through the pylorus.

NORMAL GASTRIC RESPONSE TO COMMON FOODS

On the basis of extensive studies made in the senior author's labora tory 19 the following table was constructed which contains data relative

EVACUATION TIMES AND HIGHEST TOTAL ACIDITIES FOR VARIOUS ARTICLES OF DICT

Articles of Diet (100-g portions unless otherwise stated)	Vumber of Observa toons	Highest Total Acidity (average) (ml 0 1 N alkali to neutralize 100 ml juice)	Exacuation Time (hours and minutes, average)
Beef and beef products	25	120	3 00
Bread and cereals	75	80	2 40
Cakes	29	90	3 00
Chicken*	20	125	3 15
Egg and egg combinations	90	80	2 40
Fish*	75	130	2 50
Fruits*	68	90	2 00
Gelatin* (fruit juice preparations)	5	70	2 00
Guinea hen*	2	110	4 00
Ice cream*	7	105	3 15
Ices*	4	65	2 35
Junket			2 25
Lamb and lamb products	4	65 135	3 00
Licorice .	14		3 00
Milk *	1	65	5 00
Cow	j j		
400 ml			0.00
75 ml	50	100	2 30
Human	3	45	1 15
150 ml	_		
225 ml	5	60	1 40 2 2o
Nuts* (25 to 50 g)	2	90	
Orange-albumin (2 1)	22	100	3 30 2 20
Pies	2	8o	2 20
Popcorn	29	90	1 30
Pork and pork products	3	60	3 15
Puddings	31	120	2 20
Sugars and candies	23	90	2 05
Turkey*	28	70	3 30
Veal *	2	140	9 30
(a) Market	i _	2.00	2 50
(b) "Bob	7 7	140 110	3 20
Vegetables prepared in different was a	121	75	2 15

^{*}Unjullited dats

¹⁶ See Researches and Britings Philip B Hawk and collaborators public i and distributed privately 1912

rate the glycerol from the pleces of mucous memhrane and use the glycerol extract as required in the later experiments.

PRODUCTS OF GASTRIC DIGESTION

Into the artificial gastric juice, prepared as above described, place the protein material (fibrin, coagulated egg white, or lean heef) provided for you by the instructor, add 0.4 per cent hydrochloric acid as suggested by the instructor, and keep the digestion mixture at 40° C. for two to three days. Stir frequently and keep free hydrochloric acid present in the solution as indicated by a blue color with Congored paper.

The original protein has been digested and the solution now contains the products of peptic proteolysis; i.e., denatured protein, proteoses, peptooes, etc. The insoluble residue may include nuclein and other substances. Filter the digestion mixture, and after testing for free hydrochloric acid neutralize the filtrate with sodium hydroxide solution. If any of the denatured protein is still untransformed into proteoses, it will precipitate upon neutralization. If any precipitate forms, heat the mixture to holling and filter. If no precipitate forms, proceed without filtering.

We now have a solution containing a mixture consisting principally of proteoses and peptones. Separate and identify the proteoses and peptones according to the directions given on p. 197.

GENERAL EXPERIMENTS ON GASTRIC DIGESTION

- 1. Conditions Essential for the Action of Pepsin. Prepare four test tubes as follows:
 - (a) Five ml. of pepsin solution.
 - (b) Five ml. of 0.4 per cent hydrochloric acid.
 - (c) Five mt. of pepsin hydrochloric acid solution."
- (d) Two or 3 ml. of pepsin solution and 2 to 3 ml. of 0.5 per cent sodium carbonate solution.

Into each tube introduce a small plece of fibrin and place tubes in the incubator or water bath at 40° C. for one-half hour, carefully noting any changes which occur; (Carmine-fibrin" may be used to advantage in this and the following tests under Castric Digestion. In this case, however, the experiments should be conducted at room temperature.) Now comblor the contents of tubes (a) and (b) and see if any further change occurs after standing at 40° C. for 15 to 20 minutes, Explain the results obtained from these five experiments.

2. Influence of Different Temperatures. In each of five test tubes place 5 ml. of peptin-hydrochioric acid solution. Immerse one tube in ice water, keep a second tube at room temperature, place a third in the incubator or water bath at 30°C, and keep the fourth at 50°C. Boil the contents of the fifth tube for a few moments; then cool and also keep it at 30°C. into each tube intro-

^{30 75} per cent commercial pepsin in 0 4 per cent HCl may be used.

n Digestion of firm an expension of a per cent HCI may be used.
n Digestion of firm an expension directions east obtains us indicated first by a recline of the protein due to the action of the acid and later by a disintegration and solution of the fitten due to the action of the oppin hydrochleries and. If uncertain at any time whether dispersion has kiten place, the primary has been placed by the protein of the fitten and position and the protein of the fitten of positions of the fitten of position which would indicate that dispersion has taken the objection for the fitten of the fitten

²¹ See Appendig.

duce a small piece of fibrin and note the progress of digestion. In which of the tubes does the most rapid digestion occur? Explain this.

3. The Most Favorable Acidity. Prepare three tubes as follows:

(a) 3 ml. of pepsin solution + 3 ml. of 0.4 per cent HCl. (Acidity about 0.2 per cent HCl or pli 1.3.)

(b) 3 ml. of pepsin solution + 1 ml. of 0.4 per cent HCl + 2 ml. of water. (Acidity about 0.067 per cent HCl or pH 1.8.)

(c) 3 ml. of pepsin solution + 4 drops or 0.2 ml. of 0.4 per cent HCl + 3 ml. of water. (Acidity about 0.013 per cent HCl or pl1 2.5.)

Introduce a smail piece of fibrin into each tube, keep them at 40° C., and note the progress of digestion. In which acidity does pepsin act best on fibrin? The acid decreases during digestion due to its combination with protein, so that as a determination of optimum pH this procedure is not exact. The optimum for pepsin is about pH 1.8 under the conditions of this experiment,

- 4. Differentiation Between Pepsin and Pepsinogen. Prepare five tubes as follows:
 - (a) Few drops of giveerol extract of pensinogen + 2 to 3 ml, of water.
- (b) Few drops of glycerol extract of pepsinogen + 5 ml. of 0.2 per cent hydrochloric acid.
- (c) Few drops of giycerol extract of pepsinogen + 5 ml. of 0.5 per cent sodium carbonate,
- (d) Two or 3 ml, of pepsin solution + 2 to 3 ml. of i per cent sodium carbonate.
- (e) Few drops of giycerol extract of pepsinogen + 5 ml. of 1 per cent sodium carbonate.

Add a small piece of fibrin to the contents of each tube, keep the five tubes at 40° C. for one-balf hour, and observe any changes which may have octured. To (a) add an equal volume of 0.4 per cent bydrochloric acid, neutralize (c), (d), and (e) with bydrochloric acid, and add an equal volume of 0.4 per cent hydrochloric acid. Place tbese tubes at 40° C. again and note any further changes which may occur. What contrast do you find in the results from the last three tubes? On the basis of these tests, what is the relative resistance of pepsin and pepsinogen to aikalies?

5. Comparative Digestive Power of Pepsin with Different Acids. Prepare a series of six test tubes each containing 5 mi. of a solution of one of the following acids" each solution having the same acidity of pH 2: (1) liGl. (2) lisSO₁, (3) liPO₁, (4) oxalic acid, (5) lactic acid, (6) acetic acid. Add 2 ml. of 0.5 per ent pepsin solution or 1 ml. of the gly-corol extract of the log's stomach and a small plece of fibrin. Put in a water bath and keep at 40° C. Note the progress of digestion. Can you confirm the findings of other observers¹¹ that it is the hydrogen-ion concentration (pH), rather than the nature of the acid is the school of the second of the second of the second of the controlling factor in the influence of acids on peptic digestion? Titrate 10 mi. of each of these acids (factic and acetic acids only 1 ml.)

Northrop (J. Gen. Physiol., 1, 607 (1919); 5, 263 (1922)) finds, however, that acctic acid is slightly less effective than the others apparently because of some action of the acetic acid

not on the enzyme but on the protein.

²³ IIC1 0 037 per cent, II₂SO₄ 0 059 per cent, II₂PO₄ 0 2 per cent, oxahe acid crystals 0 15 per cent, lactic and 7.2 per cent, and acetic acid 18 per cent. These solutions are best checked for pill by adding a few drops of thy mol blue solution to 5 m of each in test tubes and if necessary, adding more acid to get the same color as obtained with the oxahe acid, which can be weighed out accurately.

with 0.1 N NaOH, using phenoiphthalein as an indicator. Calculate the normality of each. Six students may conveniently work together on this part of the experiment, each student then assembling the entire data. What does this experiment teach as to the relationship of hydrogen-ion concentration rottratable acidity and as to the holosical significance of each?

6. Quantitative Determination of Peptic Activity. See Chapter 15, Gastric Analysis.

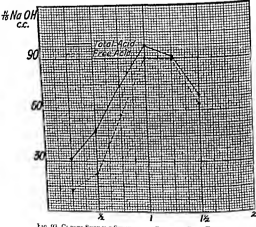


Fig. 93. Curves Showing Stimulatory Power of Beef Extract From unpublished data collected in the sensor author's laboratory by Dr. Chester C. Fowler

- 7. Quantitative Determination of Rennin. Prepare a standard mlik of pil 50 by mixing equal volumes of fresh milk and M acetate buffer of pil 50 lintroduce 10 ml portions of this milk into a series of test tubes and keep at 20° C. Add 1-ml. portions of various dilutions of a rennin solution. Mis and note the time of clotting. The amount of rennin that clots 1 ml. of the huffered milk in 10 minutes at 20° C is called one unit of rennin.
- Characteristics of Human Gastric Juice. Take some of the human gastric juice collected as described on p 369 and show that it is acid in reaction, that it contains chlorides, and that it has the power to digest protein material and to curdle mile.

^{**} Any good commercial rennet may be used in preparing this solution.

9 Chemical and Psychical Stimulation of Gastric Secretion. Have one or more volunteers from the class swallow the Rebfuss stomach tube as directed on p 384. The subjects must omit breakfast if the tube is taken in the morning or luncheon if taken in the afternoon. Empty the stomach (see pp 384 385) and, with the tube still in position, allow each subject to drink 250 ml of

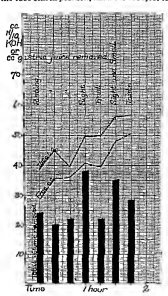


FIG 94 CURVES SHOWING PSYCHICAL STIMU LATION OF GASTRIC SECRETION Miller Bergeim Rehfuss and Hawk Am J Physiol 52 1 (1970)

bouillon prepared by dissoiving one bouillon cube in hot water Collect samples of gastric contents at intervals until the stomach is empty as described under Section 5 on p 386. The samples thus collected may be examined qualitatively for acid, chiorides pepsin, and rennin, or they may be submitted to the quantitative procedure given on p 386. If the examination is made quantitative the data may be recorded in the form of a curve such as shown in Fig. 93.

For the psychical stimulation, empty the stomach as above Then instead of having the subjects drink bouillon, simply permit them to see and smell

an appetizing beefsteak while It is being cooked. Collect samples of gastric contents as above, plot a curve and compare with Fig. 94.

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15

Gastric Analysis

The method of gastric analysis which was in vogue clinically for years entailed the feeding of a standard test meal, the removal of the complete stomach contents at the end of a one-hour period, and the analysis of the material so removed. That this method is inaccurate has been repeatedly demonstrated in the senior author's laboratory and elsewhere Furthermore, owing to the bulk of the old form of stomach tube and the discomfort occasioned by its use, it is impossible to follow the whole cyclo of digestion and estimate, step by step, the exact changes which take place in the stomach after the introduction of definite food mixtures into that organ.

Realizing the inadequacy of the procedure entailed in the old method of gastric analysis, an improved procedure was developed by Dr. Martin E. Rehfuss in the senior author's laboratory This so-called Fractional Method entails the analysis of samples of material withdrawn from the stomach (by syringe) at sbort intervals for a period of two hours or more (until the stomach is empty) after the ingestion of the test meal By this means the observer is able to follow the entire cycle of gastric digestion and is not limited, as in the old method, to information derived from the analysis of a single sample of stomach contents withdrawn at the end of one bour. That the acid values obtained by the old method may be grossly misinterpreted and lead to an incorrect diagnosis is indicated by

the diagram shown in Fig. 95.

It is set forth in Fig. 95 that various types of abnormal gastric secretion would be considered normal on the basis of a single examination at the end of one hour, whereas the application of the fractional method reveals an abnormality of the secretion. The removal of samples of gastric contents at short intervals, for a period of two hours or more after a test meal, is made possible by the use of a modified stomach tube of small diameter (No. 12 French tubing) and fitted with a metal tip. The tip is slotted with large perforations, the diameter of each being equivalent to the maximum bore of the tubing Such a tube can be left in the stomach through the entire cycle of gastric digestion without inconvenience to the patient. The Rehfuss stomach tube is shown in Fig. 96.* Lyon suggested a modified tip

A tube much favored in England is that devised by Ryle.3 This consists

Rehfuss Am. J. Med. Sci., 147, S48 (1914).
 Thus tube is manufactured by Charles Lentz and Sons, Philadelphia.

^{*} Byle, Gastric Function in Health and Disease, London, Oxford University Press, 1926.

of a small bore rubber tube with a blind end, into which is inserted an oval weight of lead. Holes are punched in the rubber tube just above the weight.

Other tubes or plastic catheters have been employed in gastric analysis ⁴ Certain of these (Levin) may be passed through the nose Such tubes are

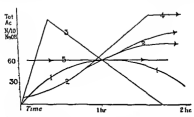


FIG 95 NORMAL AND PATHOLOGICAL CURVES AFTER AN EWALD

(1) Normal curve (2) delayed digestion with late hyper acidity (3) larval hyperacidity, (4) tardive hyperacidity and (5) marked continued secretion from obstruction



FIG 96 REBFUSS STOMACH TUBE

useful in rapidly demonstrating the absence or presence of free hydrochloric acid in the stomach contents

A method of determining gastric acidity without initiation has also been suggested by Segal and his associates * An editorial in the Journal of the American Medical Association* discusses the method as follows

One of these the Sawyer tube was developed at the Mayo Clinic by Miss Catherine

Seral Miller and Morton Proc Soc Expli Biol Med 74 218 (1950) Segal Viller Morton and Young Gastroenterology 16 380 (1950) Segal Vied Clin N Amer 35 593

J Am Ved Assoc 146 260 (1951)

'Segal and bis associates recently described a test for determining the presence or absence of free hydroclibric and in the stomach witbout subjecting the patient to intubation. The principle if the test hes in the use if a cation exchange resin to which is attached a special indicator cation that can be readily identified when it is released from the compound by an inn exchange reaction. Clinical trials have been based in administration of a quinnium exchange indicator compound' prepared by subjecting an acid-conditioned cation exchange resin that a solution of quinnine hydrochloride. By this treatment quinnine replaces the bydrogen cations of the carboxylic acid groups present in the resin. If free hydrochloric nead is present in the stomach the quinnine will be absorbed into the body and can later be detected in the urine. Smaller amounts of quinnium cations may be displaced by the cations present in the secretions of the small intestine. However, it has experience in the stomach the smaller statine However these quinnium cations can be differentiated from those displaced by the hydrogen ions in gastric juice by the time of appearance and the amount of quinnium cations in the urine.

This procedure is more pleasant for the patient than the conventional intubation method. Its simplicity suggests its usefulness as a screening test to select achiorhy drie subjects in the gastric cancer age group fur further investigative studies?

PROPERTIES AND COMPOSITION OF HUMAN GASTRIC JUICE

	Appetite Juice	Residuum
Specific gravity	1 007	1 006
Freezing point depression °C	-0 55°	-0 47°
Total acidity, per cent HCl	0 45	0 30
Total solids g per 100 ml	0 55	0 98
Organic solids g per 100 ml	0 41	0 53
Inorganic solids g per 100 ml	0 14	0 45
Total nitrogen g per 100 ml	0 060	0 066
Total phosphorus g per 100 ml		0 005
Total sulfur g per 100 ml)	0 007
Ammonia N. g per 100 ml	0 002 3	
Amino acid N. g per 100 ml	0 003 0	
Chlorides (as Cl) g per 100 ml	0.5	

An apparatus has also been devised for the determination of intragastric conductance and temperature. The apparatus is also provided with an aspiration tube similar to that of the Rehfuss tube which makes possible the removal of samples of gastric contents for chemical analysis.

For a long time the consensus, based principally upon the work of the Pavlov school, was to the effect that the gastric juice of normal man had an average acid concentration of 0.2 per cent hydrochloric acid, whereas the gastric juice of the dog and cat had an inverage acid concentration of 0.4 to 0.5 per cent hydrochloric acid. These experiments were based principally upon the examination of the pure gastric juice of the lower nimials as compared with the stomach contents of man. Later experiments, however, demonstrated that the acid concentration of the freshly

⁷ Diagnex (Squibb) is useful in this connection. The appearance of quinine in the urine within two hours after the administration of Diagnex aboves the 1 resence of IICl. The absence of quinine within this period indicates achievely dria.
3 Bergaim Am J Phusid 45.1 (1912).

secreted gastrie pines of man is similar to that of the dog 1 e 0 4 to 0 5 per eent hydrochloric acid. Boldvreff claimed that this initial high acidit of the human gastrie pines is normally lowered to the "optimum acidity" of 0 15 to 0 2 per cent by reguritation of alkaline fluids (bile pancreatic and intestinal piness) from the intestine. This constitutes what Boldyreff termed 'the automatic regulation of gastrie acidity." This claim has been amply substantiated. Both bile and trypsin are easily identified in the stomach contents of man after the introduction of 0.5 per cent hydrochloric acid into the empty organ.

The composition of human gastric junee and of the residuum (see p 384)

is given in the table on p 377

THE USE OF INDICATORS IN GASTRIC ANALYSIS

Indicators are used in gastric analysis for two main purposes. (1) For the colorimetric determination of the plI of gastric contents and (2) as an aid in the measurement of gastric heidity by titration. The use of indicators for the first purpose is identical in principle with their use in this connection with other fluids and is described in detail on p. 37. The practical details in connection with gastric analysis are presented on p. 379.

The primary objective in the titration of gastric acidity is to determine the amount of unneutralized hydrochloric acid present (free hydrochloric acid') in the possible presence of other acids which while titrata ble are nevertheless so much less highly ionized than hydrochloric acid that they contribute little or nothing to the hydrogen ion concentration of the solution It is the hydrogen ion concentration (i.e. the pH) of the gastric contents which to a large extent determines whether or not the physiological functions of the gastric secretion will be served and hydrogen ions in concentration sufficient to maintain a normal pH caa come only from a highly ionized acid such as hydrochloric acid Thus gastric function can be evaluated in terms of the presence and amount of free hydrochloric acid Since the concentration of free hydrochloric acid determines the pH it is clear that a pH measurement will frequently give as much information as a titration and increasing use of pH determina tion in this connection is being made elimically particularly in view of the ease with which precise pH determinations may be made with the modern electronic pH meters (see Chapter 1)

The establishment by titration of the presence and amount of free hydrochloric acid in the presence of o her titratable acids is based on the fact that hydrochloric acid is completely dissociated in solution the hydrochloric acid is completely dissociated in solution the hydrogen ions from this dissociation reacting with the added OH ions before any undissociated acid present can ionize and so react. Thus the amount of alkali added up to the point of practically complete neutralization of the hydrochloric acid present should be distinguishable from that necessary for the remaining acid or acids. That this is so is cyident from an inspection of the titration curve for hydrochloric acid as compared with that for a typical weak acid such as acetic acid such curves are shown in Fig. 11 of Chapter 1. From an inspection of these curves it can be seen that as standard alkali is added to a solution containing hydrochloric

acid, the pH of the solution changes relatively little until most of the acid is neutralized, increasing from pH 1 to about pH 2, when about three-fourths of the acid has been neutralized. As the titration continues, at nbout pH 3 5 or so it is clear that practically all of the acid has been titrated, stopping the titration at this point gives a value for the acid present which is almost indistinguishable from the value obtained if the titration were carried to the end point with such indicators as methyl orange, litinus, or phenolphthalein

The situation is quite different if a weak acid, such as acetic acid for example, is being titrated. In the case of acetic acid, the solution has pH of about 3 before any alkah has been added, this pH corresponding to the relatively small (about I per cent) ionization of the acetic acid molecules. As the pH is increased by the addition of alkah, more of the acetic acid dissociates to give hydrogen ions which are capible of reacting with the added OH ions until ultimately sufficient alkah has been added to neutralize all of the acid initially present. But it will be noted that the end point of this titration (i.e., the pH at which equivalent amounts of alkah and acid are present) is not pH 7 but rather nearer pH 8.5. Thus, in order to titrate such an acid as this, it is necessary to use an indicator which changes color at about pH 8.5, phenolphtbalem is such an indicator.

From what has just been said, it follows that if a mixture of hydrochloric acid and some weak acid or acids is being titrated with standard alkali, the buret reading at pH 35 or thereabouts will be a measure of the hydrochloric acid present, while the reading at pH 8 5 will be a measure of the total acidity of the solution. It is thus possible to distinguish quantitatively between these two types of acidity provided that means of indicating the pH of the solution are available. This may be done using a pH meter, it is much more common to select an indicator whose color change hes at the pH range desired Of the various indicators which have been proposed for this purpose in gastric analysis, Topfer's reagent (dimethylaminoazobenzene) and phenolphthalem are almost universally used Topfer's reagent has a color change from red to yellow over the pH range 2 9 to 4 0 (see Table), the intermediate color of salmon pink being noticeable at approximately pH 3 3 Thus if gastric contents are titrated with alkali to the color change with Topfer's reagent, a measure of the free hydrochloric acid present will be obtained, the value being uninfluenced by any weak reads which may be present. If the titration is then continued to the color change with phenolphthalem (pH 85), the total acidity is determinable

Topfer's reagent has a number of disadvantages, the color change is not sharp and requires a certain amount of familianty before the proper end point is routinely obtainable, furthermore the color fades rapidly at the end point and thus precludes the setting up of pH controls for the more precise establishment of the end point Other indicators have been proposed, such as thymol blue (red, becoming yellow at pH 2 8), and bromophenol blue (yellow, beginning to turn blue at pH 3 4 or so), but in the authors' experience of teaching medical students and technicians the routine of gastric titrations, neither of these has proved so satisfactory as Topfer's reagent.

In the above discussion acctic acid was used as an example of a typical weak acid to illustrate the principles involved in a gastric titration. In actual practice the weak acids which may be found in gastric contents include protein hydrochloride (so-called "combined by drochloric acid'), acid phosphates and various organic acids such as lactic, citric etc after fermentation or the ingestion of certain foods. At one time the mistaken notion prevailed that by the suitable use of various indicators it was possible to differentiate between these components of the weak acid frac tion of gastrie contents. This is not true since the titration curves of these various components overlap to such an extent that it is impossible to differentiate between the contribution of each to the total acidity, and the concept should be ahandoned Even the distinction between free mineral acid and weak organic acids becomes less sharp if the organic acids have an appreciable ionization at pll 3 or so, as is the ease for example with lactic acid. The presence of significant amounts of such organic acid in gastric contents is quito unusual, should it occur the determination of volatile chloride is of value in establishing the extent of acidity due to hydrochloric acid

It is occasional practice in the titration of relatively pure gastric contents to subtract the value for the free acidity from that of the total acidity and call the difference the "combined acid." The valuity of this is questionable even assuming that the difference between free and total acid is partly due to acid which has reacted with protein to form protein hydroehloride, it is clear that the amount of acid which has so reacted is measurable only by titrating the solution to the isoelectric point of the protein, further titration beyond this pH represents the formation of alkah salt of protein and will depend on the amount and nature of protein present without reference to how much hydrochloric acid has been "combined" with the protein Actually the difference between free and total acid is more a measure of the huffer power of the gastric juice than any thing else From a practical point of view therefore it would appear that the requirements of gastric analysis at least in so far as gastric acid is concerned, are largely met by measurement of either the free acid or the pH

The table on p 381 lists the characteristics of those indicators which have found application in gastric analysis and for other purposes Experiments which follow illustrate the application of the principles just presented

Tests with Indicators Prepare a series of solutions of varying acidities as outlined in the following table, p. 332 introduce 5 or 10 ml portions of each of these into a series of test tubes and to each add a few drops of a solution of thymol blue. Make a note of the colors produced, in the spaces jeft for this purpose in the same way test the other indicators mentioned, in order, in each case using a few drops of the indicator solution.

Are the following assumptions on which the use of certain of these indicators in gastric analysis is based borne out by your findings?

1 That Topfer's reagent (dimethylaminoazobenzene) gives its characteristic pinkish red color only in the presence of free HCl

2 That Congo red can be used to distinguish between strong acids, moderately weak acids, and very weak acids

moderately weak acids, and very weak acids

3 That thymol blue may be used as an indicator in the titration of

botb free and total acid

4 That alizarm may he used in titrations where the end point is at
a pH just acid to pH 7 rather than just beyond pH 7

5 That phenolphthalem can be used in titrating total acidity, that is,

acidity due to mineral and organic acids, acid sults, and combined acid
6 That Gunzberg's test is in certain respects the most satisfactory

one for free HCl
7 That "combined acid" (protem hydrochloride) is an acid of approximately, the strength of acetic acid

	pH Range	Color Change
Thy mol blue (read range)	1228	Red yellow
Topfer s reagent	29-40	Red yellow
Bromophenol blue	30-46	Yellow blue
Congo red	30-50	Blue-red
Methyl orange	3 1- 4 4	Orange red yellow
Bromocresol green	40-56	Yellow blue
Methyl red	4 2 6 3	Red yellon
Litmus	45~83	Red blue
Chlorophenol red	60-66	Yellow red
Alizarin red	60-68	Yellow red
p-Nitrophenol	60-70	Colorless yellow
Bromocresol purple	5 4- 7 0	Yellon purple
Bromothymol blue	60-76	Yellon blue
Phenol red	66-82	Yellow red
Neutral red	6880	Red yellow
Cresol red	7 2-88	Yellow red
Meta cresol purple	76-92	Yellon purple
Thymol blue (alkaline range)	8 2- 9 8	lellow blue
Phenolphthalein	8 3 10 0	Colorless red
Alizarin yellow	10 0-12 0	Colorless vellow
Tropaeolin O	11 1 12 7	I ellow-orange

Special Test for Free HCl (Gunzberg's Test) Perform the following test on Solutions 1 to 4 of the table and tabulate the results Place 1 to 2 drops of Gunzberg's reagent' in a small porcelain evaporating dish and carefully evaporate to dryness over a low flame. Insert a glass stirring rod into the mixture to be tested and draw the moist end of the rod through the dried reagent Warm again gently and note the production of a purplish red color in the presence of free hydrochloric acid. This test differs markedly from the use of indicators in that the reaction is not determined by the pill of the solution, but is based on the fact that hydrochloric acid of any strength whatever reaches a "constant boiling" concentration of about 20 per cent on evaporation. At this strength of acid the ingredients of the reagent condense to form a purplish red compound. The reaction is thus highly specific for hydrochloric acid in gastric fulce.

See Appendix

TABULATION OF REQUESS OF TESTS ON INDICATORS.

		TVI	חדיי			TARGENTING		,	,	•	6	10	=	13	2
	-		-	•	*5	4	•						101		
	- roug			787	Vethal	Вгото	Congo	Bromo-	Chlorophenol	\$ 5	Bromo	Cresol	phthal	Ganz- berg's	
į	701	_	To M		Orange		Red	Green	Red	e C	Blue		Ē	Reagent	
Solution	mality or	nate 114	antr	Venger.	:	30-50 50 66 50-68	30-50	40-56	50.66	50-08	60-76 7288	7288	8.3-100	341)	
	Votar		8 2 2 2 8	2.0	1		-								P
									i			4			RA.
	-	:			_										CTI
I, 0.4 per cent HC1	N/10	2													CA
2, 0 04 per reat 11C?	N/100	7 0													LI
	N/10	000				_							-		11.3
3. 0.0 per cent acette acit		-		Ī	1										151
4 0 04 per cent combined 11Cit	N/100	0							T	1	I				ULA
5 Acid phonibate 9 14	2 10	0 0					Ì								ж
0 Acid histinate basic lice liste 4 01	1 1/1	0,						Ì							AL C
7 Barie lice; have 20 1\$	M/13	8 0			j		Ì	Ì							12.81
8. Borate \a0il 6 41	N/10	10 0				Ì		Ì			-				STR
9 04 prent Vall	N/10	13 0			_										
# In Insternations, Thund Bir Creater Bromophron May Bromerred green Chlorophrond red, and Bromothymod Mac 0.01 g. in 100 ml of alreado Verled overnor OI g in	Cresol red	Bromophe	Inol Mus L	Promecres	d green G.	Morepheno	t red, and	Bromothy	not blue 0	01 K in l	30 mt of a	leohol V	ethyl orang	# 01g ln	

or Combined 11CL. Treat 0 4 per cent 11Cl with a small amount of Witter s peptons and bod until the solution no longer gives a blue but only a brown color with Congo-red 100 ml. ol nater Topfer's reggent, 0 5 g. el dimetly lammonandementana in 100 ml of 05 per centalound. Compo red, 0 3 g in 90 ml of nater and add 10 ml of her centalentol. Alteren, In of solium alianto sulfante in 100 ml of water Phenolystholem, In in 100 mt of 93 per cent alredtol

this our solutions of potassium dity drogen those they had a desedum hydrogen phosphats of M/18 strength See 35 To prepare the acid; host hate solution used in the feet wit I part of the solution of the discolumn selt with 9 parts of the solution of the abid drosers I keet hat I ber the I have planted as colution the proportions are 20 1, for

| Rente-NaOII solution Pres are a bornte solution by dissoluing 12 404 gt of pure 1 rio at [10 2 mai] in 100 ml of N NaOII solution and oblite whith water to a liter Presthe solution of 1117, the 1 reportions are 6 4 Do not attempt to use destined 1140 alone as a pil 7 solution (Why not?) pare the burate-NaOII solution by mixing 6 parts of the lorate solution with 4 parts of 0.1 N NaOII

Differential Titration of a Mixture of Strong and Weak Acids (a) Titrate a 5-ml portion of 0 1 N acetic acid with 0 1 Nsodlum hydroxide, using phenolphthalein as indicator (b) Titrate a 5-mi portion of 0 1 N hydrochloric acid. using phenolphthalein (c) Titrate a 5-ml portion of 0 1 N hy drochloric acid, using Topfer's reagent (one drop) as indicator. The end point is a salmonnink shade, intermediate between red and vellow. Is there any significant difference in the titer of the hydrochloric acid as compared with the value ohtained in (b)? (d) Now mix 5-mi portions of 0 1 N acetic acid and 0 1 N hydrochloric acid in a flask, add Topfer's reagent, and titrate with alkali to the salmon-pink end point. Read the huret, add a drop of phenolohthalein to the contents of the flask, and continue the titration with alkali until the pink color of the phenolphthalein end point can he seen to be superimposed on the clear yellow color of the Topfer's reagent Read the buret again This reading represents the "total acidity" The first reading is a measure of the HCl present, and the difference between the first and final readings represents the acidity due to the weak acid (acetic, in this case) Compare the values you obtain In this differential fitration with those obtained by separate analysis in parts (a) and (c) above

THE FRACTIONAL METHOD OF GASTRIC ANALYSIS

Since 1914, when the first experiments entailing the use of the fractional method were reported from the senior author's laboratory, the method has been widely adopted hoth in this country and abroad. This wide-spread use by a large number of workers has resulted in much discussion of the method. On the basis of this large experience certain modifications in the original technique are here presented in its major details, however, the method remains as originally carried out, and in the opinion of the authors constitutes the hest available method for the clinical examination of the stomach

When an Ewald test meal is given to normal induiduals a curve such as Curve (1) io Fig. 95 is usually obtatoed. The curve may vary within certain limits depending on individual dicks increases but it is usually found to follow the curve depicted and the meal oormally leaves the stomach in two and one-half hours. Pathologically every variation occurs in time of exacustion as well as so the character of the curve and the quantity of the secretion elaborated Fig. 95 represents some of the possibilities of pathological cases but a consideration of their interpretation is outside the purpose of the present volume. It is evident however, that the cycle of gastric digestion is a constantly changing one and no information cooccuring the trend of digestion can be obtained by an examination at only a single stage of digestion. Marked changes may precede or follow that stage.

Low acidities may be found in caremona in atomic dyspepsia and in permisons menina. However, aniedity of itself is not necessarily of pathological significance. A tendency toward high acidities may be found though not constantly, in cases of gastric and duodenal ulcers especially those occurring in the oeighborhood of the pylorus and indusing some degree of obstruction. High acidities may be induced reflectly in gallibladder discusse and appendictus.

As this is written one of the favorite means used to combat the high acid tendency of ulcer is the chemical substance called Banthine (B-diethy laminoethy kanthene-9-ariboxy late methobromide). However, there are those who do not believe that Banthine is the final answer to the ulcer problem "It is suggested that the drug

¹⁸ Benjamin Rosiere and Grossman Gastroenterology 15 727 (1950)
¹⁸ Ruffin Gastroenterology 17 559 (1951) Levin Kursner and Palmer Gastroenterology
21, 339 (1951)

should be used to supplement conventional methods of treatment rather than to replace them. Pro banthine and Prantal have also been recommended for use in user therapy. In the past rocation therapy as well as the use of radioactive phosphorus (P2) and radioactive indine (I¹¹) have been used with indifferent success to suppress gastrie acidity clinically Radioactive krypton may possibly prove more effective for this purpose ¹¹ However this has not been definitely praved

Some gastroenterologists have found the gastrophotor useful in the study of uler. This is a tiny camera which may be swallowed by the patient. By its use 16 pictures of

various sections of the stomach walls may be taken at the same time

OBTAINING THE SAMPLES

I Introduction of the Stomach Tube Whereas the large tube is directly inserted by propulsion, the Rehfuss tube is swallowed in the natural manner with the aid of gravity The tube may be passed in one of three ways, viz, (1) lubricated, (2) with the aid of fluid, (3) after the throat is cocainized When passed by the first method the tip of the tube, after thorough lubrica tion with glycerol or liquid petrolatum is held between the thumb and fore finger and placed on the tongue Then with the aid of the forefinger the tip is pushed backward until it reaches the root of the tongue and is engaged in the oropharynx Then the patient is encouraged to breathe deeply through the nose and to swallow persistently while the tube is slowly fed into the mouth After slight discomfort in the pharynx and its passage past the level of the cricold cartilage, practically no discomfort is felt. This method is used when it is essential that the pure gastric secretion or residuum be obtained Ordi narrly, however, it is much easier to swallow the tube by the second method This method consists in placing the tip in the oropharynx and then giring the patient a measured quantity of water or tea to swallow The movements induced by the swallowing carry the tube rapidly to the stomach with a minimum of discomfort When as Ewald meal (ses p 385) Is given, part of the tes may be reserved for swallowing the tube This procedure makes it scarcely more arduous than the swallowing of food Should the patient, however, be extremely neurotic or the unfortunate possessor of marked pharyngeal byperesthesia, cocaine hydrochloride in 2 per cent aqueous solution may be applied to the throat, rendering the passage of the tube practically insensible When the tube has entered the stomach, aspiration of the material shows the characteristic gastric contents Should the tlp remain in the esophagus through transient cardiospasm or other cause, aspiration results in the re moval of only a very small specimen having all the characteristics of the pharyngeal and esophageal secretions

The clinician should see to it that the tube is so placed that the tip reaches the lower pole of the stomach Furthermore, the tip should remain in this position throughout the duration of the test By placing the tube in this manner, representative specimens and accurate data may be obtained

It has been shown that the gastric response to the fractional method when properly performed is similar for the same individual on different days."

Bioomfield and Keefer have suggested a method for the "continuous quantitative estimation of gastric secretion and discharge in man"

2 Removaf and Analysis of Residuum II the so called empty stomach is examined in the morning before any food or drink has been taken it will be found to contain considerable maternal, which is termed rendum Before's

¹⁹ Steinfield Proc Soc Papil Biol Med 81 636 (1952)

¹¹ Ryle Gastric Function in Health and Disease London Oxford University Press 1976

test meal is introduced into the stomach, this organ should be emptled. If this is not done we cannot consider the samples withdrawn after the test meal is eaten as representing the secretory activity of the gastric cells under the influence of the stimulation of the test meal. It has been generally recognized, clinically, that a residuum greater than 20 ml. is pathological. Such a volume has been considered as indicative of hypersecretion, and this in turn In many cases indicates an organic lesion. The observations indicating that a residuum of over 20 ml. was pathological were made upon residuums removed by means of the old type of stomach tube which does not completely empty the stomach. When the residuum is completely removed by means of the Rehfuss tube it has been demonstrated that the normal residuum is practically always over 20 ml., the average for both men and women being about 50 ml. The normal residuum has been found to possess all the qualities of a physiologically active gastric juice with an average total acidity of 30 and an average free acidity of 18.5. The residuum is often colored by blie. This is particularly true if the fluid has a relatively high acidity. Trypsin is also generally present. These findings indicate regurgitation (see p. 390). Pathological residuums may contain blood, pus, and mucus and may also show food retention, indicative of delayed evacuation. In carcinoma the residuum frequently has a foul odor. The quantity may also be much increased owing to hypersecretion. A residuum of large volume possessing a total acidity value of 70 or over may indicate ulcer.

Remove the residuum as directed under (5) on p. 386, and onalyze the fluid according to methods outlined on p. 386.

3. Feeding the Test Meal. Before making an analysis of the stomoch conents it is customary to introduce something into the stomach which will stimulate the gastric cells. The response to this stimulation is then measured clinically by the determination of total acidity, free acidity, and pepsin in the stomach contents. Many forms of test meal have been used.

The test meal most widely employed is the Ewald test meal. This consists of 2 pieces (35 g.) of toast and 8 ounces (250 ml.) of tea.

Inasmuch as it was demonstrated in the senior outbor's laboratory that water gave a similar gastric stimulation to that produced by the Ewald meal, it was suggested that a simple water meal might be substituted for the Ewald meal. This water meal also has the added advantage of enabling one to determine the presence of food rests and to test more accurately for iactic acid, blood, and bile. Rylet' prefers to use oatmeal gruel (1 pint). Another test meal favored in certain English hospitals consists of 20 g. citrus pectin, 35 g. sucrose, 60-70 mg. phenol red and sufficient caustic soda to raise the pH to 6.5 in one liter of distilled water. The alcohol test meal has also been sugested. This coosists of 200 ml. of 5 per cent ethyl alcohol. Doses of 0.5 to 1 mg. of histamine hydrochloride injected subcutaneously are effective in stimulating gastric secretion and may help to differentiate between true and "apparent" achilla dastrica.

4. Feeding the Retention Meal. In order to obtain more information regarding gastric morility than is furnished by the ordinary test meal described above, the patient may be fed a so-called retention meal. This meal is fed in place of the regular evening meal and contains substances readily detected.

¹⁴ Bod 2 thep of catmeal in a quart of water until the total bulk is reduced to 1 pint Strain through coarse muslin. Season with salt as desired.

¹³ Hunt. Lancet, 257, 794 (1949), and J. Physiol (London), 113, 169 (1951)

¹⁶ Ehrmann. Am J. Digestire Diseases, 13, 23 (1916).

in the morning before breakfast (7 to 8 A M), remove the atomach contents (residuum see p 384) hy aspiration and examine for food rests The normal stomach should give no evidences of food retention A satisfactory retention meal consists of 4 ounces each of bolled string beans and rice Diets contain ing prunes raspberry marmalade, 13 copodium powder, etc., have also heen employed in many instances an ordinary mixed diet will serve the purpose

5 Removal of Samples of Stomach Contents At intervals of exactly Is minutes from the time the test meal is eaten until the stomach is empty, 5 to 6 ml samples of gastric contents are withdrawn from the stomach by means of aspiration (A few milliliters of air is blown down the tube in order that the tube shall be empty when the next sample is drawn) if the stomach is not empty at the end of three hours, the remaining stomach contents may be withdrawn and measured

In order to facilitate the mixing of the stomach contents and the with drawal of a more representative sample, the stomach contents should be aspirated back and forth four times before taking the sample for analysis Some clinicians advise that the stomach contents be mixed by physical manipulation just prior to asplatation

In the removal of samples from the stomach, it is essential that very little traction be employed. To completely empty the stomach, aspiration is practised in four positions (a) on the back, (b) on the stomach, (c) on the right side, and (d) on the left side. This results in complete evacuation of the stomach. Three tests may be employed to determine whether the stomach is empty (i) No more material can be aspirated in any position (2) Injection of air and auscultation over the stomach with a stethoscope reveals a stick rale and not a series of gurgling rales such as is heard when there is material in the stomach (3) Lavage or irrigation through the tube shows the absence of all food in the stomach.

EXAMINATION OF THE SAMPLES

The old methods of gastric analysis involved the collection (by analys a and calculation) of data regarding six-real types of acidity. It I as already been pointed out (9 380) that the validity of such data is questional le. The modern tendency among clinicians is to lay particular emphasis upon it evalues for total acidity. Free acidity and pill. Chloride partition may also give some information. The determination of the peptic activity is also of occasional value as well as it is demonstration of the presence or absence of occult blood facine acid mureus food resist etc.

Strain each sample through fine mesh cheesecloth ¹¹ Examine the residue for mucus blood, ¹¹ and food rests. Use the strained stomach contents for the determination of total acidity, free acidity, and peptic activity by methods which follow

Determination of Total Acidity Principle The indicator used is phenol
phthalein Since tle in heator reacts with mineral acid organic acid combined acid
and acid calls tle values obtained represent tle total acidity of tle solution

Procedure Measure I mi of the strained stomach contents by means of an Ostwaid pipet and introduce it into a low form 60 mi porceiain evaporating

*The detection of flood is rather more satisfactory in the resulus than in the strained fluid.

is Any examination required for microscopic constituents should be made on the right (units ned) gastric contents. Tests for occult blood may be made on the sediment if des red.

dish Dilute with 15 ml of distilled water Add 2 drops of a 1 per cent alcoholic solution of phenoiphthalein and titrate with 0 01 N sodium hydroxide until a finint pink color is obtained which persists for about 2 minutes Take the huret reading and calculate the total acidity

CALCULATION Note the number of milliliters of 0 01 N NaOH required to neutral ize 1 ml of stomach contents and multiply it by 10 to obtain the number of milliliters of 0 1 N N nOH necessar; to neutralize 100 ml of stomach contents This is the method of calculation most widely used

2 Determination of Free Acidity Principle An indicator is selected which changes color at sufficiently low pH values so that the end point corresponds to the neutralization of all highly ionized acid only, as discussed on p 378. The indicator most widely used is Topfer's reagent its advantages and disadvantages have been presented on p 379. The use of Sahli's reagent a mixture of potassium iodide and iodate which liberates free iodine in the presence of a sufficiently high concentration of hydrogen ions has been suggested in place of Topfer's reagent. The liberated iodine is titrated with thiosulfate solution and the end point is very sharp Unfortunately, Sahli's reagent reacts to n certain extent with weak organic acids so that high values are obtained for example after the ingestion of acid fruits. A procedure using Sahli's reagent is given in the eleventh edition of this book.

Procedure By means of an Ostwald pipet measure 1 ml of the strained stomach contents and introduce it into a low form 60-ml porcelain evaporating dish Dilute with 10 ml of distilled water Add 2 drops of Topfer's reagent (0 5 per cent alcoholic solution of dimethylaminoazobenzene) and titrate with 0 01 N sodium bydroxide to a salmon pink color if the sample gives a yellow color on the addition of Topfer's reagent, it has no free acid present The end point may require practice in identification. The use of comparative buffered colorimetric standards to control the end point has been suggested 10 Care should be taken against the possibility of color fading. The reading of the buret at the end point is a measure of the free acidity.

It is common clinical practice to use this same sample for the determination of total acidity. To do this, add 2 drops of a 1 per cent alcoholic solution of phenolphthalein, and continue the titration until the pink color change of the phenolphthalein is superimposed on the yellow color of the Topfer's reagent The buret reading at this point is a measure of the total acidity.

CALCULATION Note the number of millithers of 0.01 N sodium hydroxide required to reach the end point color with Topfer's reagent Multiply this by 10 to obtain the number of millithers of 0.1 N sodium hydroxide necessary to neutralize the free acid in 100 ml of gastric contents. This value also corresponds to the free neudity expressed in degrees or in millicquivalents of acid per liter. In a similar way, the buret reading at the phenolphthalein end point can be used to calculate the total acidity.

3 Determination of Hydrogen ion Concentration of Gastrie Contents Principle Inasmuch as the hydrogen ion concentration of the gestric contents as a determining factor in peptic digestion the estimation of pH is coming into wider use in gastric analysis. Simple colorimetric methods are an adiable for this purpose, as is the concenient electrometric pH meter. The measurement of pH gives information of real value and the test paper method especially is so convenient that it may often be the method of choice. Determinations of pH do not, however, displace titration methods, since the latter give additional information.

¹³ Berk Thomas and Rehfuss Am J D gestire Diseases 9 106 (1942)

²⁸ Prepared outfits for both the indicator solution and test paper methods may be obtained from nost larger laborator; supply houses

(a) COLORIMETRIC METHOD OF SHORL AND KING

Procedure Prepare Clark and Lubs standard solutions for pH 14, 16, 18 20 24, 30 (see Chapter 1) Transfer 2 ml of filtered or centrifuged gastric contents to a test tube 11 mm in diameter Add 2 drops or 0.04 ml of 0.2 per cent thymolsulfonephthalein in 95 per cent alcohol Compare with equal amounts of standard solutions treated in the same way in similar test tubes if the sample is more acid than pH 14, lower standards may be prepared Using the Ewald test meal it may be said in a general way that pH 14 represents a high acidity, from 16 to 18, moderate normal acidity, from 20 to 24, low acidity, and 30 or higher, anacidity

(b) METHOD OF DENIS AND SHEVERMAN USING TEST PAPERS

Procedure Prepare test papers as follows A solution of dye (dimethyl aminoaxobenzene 0 5 per cent or 0 1 per cent thymol blue) is poured over a sheet of filter paper six inches in diameter, the excess dye allowed to drain of, and the paper dried quickly by holding it in front of a rapidly revolving electric fan Prepare the two kinds of paper, cut in strips approximately 1 by 1/2 inch, and place in a stoppered bottle Keep away from light and laboratory fumes Smooth, ashless quantitative paper of the highest grade should be used

Pour some of the material to be tested over the end of a test paper. The colors may be compared with those obtained from similar papers dipped in the standard solutions mentioned in the method of Shobi and King above However, with some practice this will be unnecessary. All readings should be made at once. The table given below will serve as a guide in the interpretation of results it gives test paper results as compared with number of mill of 0 l. N NaOII that would be required to neutraine 100 mill of a solution of the ii ion concentration found, if all the acid were free iiCl

Indicator	pii	Color	Ml 01 N NaOH Required	Remarks
Topfer's reagent Topfer's reagent Topfer's reagent Töpfer's reagent Thymol blue Thymol blue Thymol blue	1 4 1 6 1 8 2 0 2 0 2 4 3 0	Deep red Reddish-orange Orange Yellowish-orange Purple red Faint pink Yellow	44 27 17 10 10 4	Hyperacidity Normal acidity Normal acidity Hypoacidity Hypoacidity Hypoacidity Anacidity

4 Determination of Peptic Activity

(a) Photometric Merindo or Ricos and Stadie. In this method the enzyme activity is measured photometrically as the decrease in turbidity of a stand ardized, homogenized suspension of coagulated egg white under specified conditions. Accurate measurement is possible down to a level equivalent to 1 kg of crystalline pepain Protein hydrolysis so measured follows a non-molecular course and hence the enzyme activity is expressed as a velocity constant. This method is particularly adapted to clinical studies. For details of procedure see the original article?

^{11] 1}ggs and Stade J Biol Chem 150 463 (1943)

(b) HEMOGLOBIA METHOD OF ANSON AND MIRSKY PRINCIPLE The pepsin is added to a hemoglobin solution. Unaltered protein is removed with trichloroacetic acid. The amount of digestion products is determined colorimetrically using phenol reagent which reacts with tyrosine tryptophan and cysteine groups tyrosine being used as a standard.

For a discussion of the determination of pepsin by viscosimetric and other methods and of pepsin units see Northrop 22

Procedure Pipet 5 ml of a 2 per cent solution of dialyzed ox carbon monoxide hemoglobin²¹ in 0 06 N HCl into a 175 by 20 mm test tube and bring to 35 5° or 25° C Add 1 ml of enzyme solution and mix by whirling tube After 5 minutes ndd 10 ml of 4 per cent trichloroacetic acid from another test tube and pour back and forth to mix Filter through fine paper To 3 ml of filtrate in a 50 ml Erlenmeyer flask add 20 ml of water, 1 ml of 3 85 N NaOH, and 1 ml of phenol reagent (Folin and Giocalteu See Chapter 31) The standard consists of 3 ml of 0 1 N HCl continuing 0 15 mg of tyrosine (A copper sulfate standard can be used with a red color filter) Compare colors after 5 to 10 minutes with the standard at 20

CALCULATION 24 If A 18 the reading of the unknown

Pepsin Units =
$$(0.0194/\lambda) - 0.000147$$

If carried out at 25° C multiply by 1 82

(e) METHOD OF VOLHARD AND LOHLEIN PRINCIPLE The pepsin is added to case in solution. The unaltered casem is salted out. The filtrate from this contains the digestion products of casem which can be estimated by titration.

Procedure Into each of three flashs (graduated at 300 and at 400 ml) introduce 11 ml of N HCl and add water to make not quite 150 ml With con-

The tyrosine used should have been recrystallized and the concentration also determined

by the Kieldahl method N = 774 per cent

14 In addition to the 0.15 mg of tyrosuse the standard contains from the reagents an amount of color producing naterial equivalent to 0.015 mg of tyrosuse Tle color producing substance in the 3 ml of filtrate is therefore equivalent to 20.024, 0.015 + 0.015) mg of tyrosuse Ol this 0.015 mg are due to the color producing substance in the reagents and 0.01 mg to the color producing substance present in the trichloroactic acid filtrate even when no enzyme is added. The digested hemoglobin in the 3 ml of filtrate is therefore equivalent to (20.7%) (0.15 + 0.015) - 0.015 - 0.01 mg of tyrosuse. This value must be multiplied by 1% to obtain the digested hemoglobin in the whole 16 ml of filtrate instead of the 3 ml taken for analysis it must be divisted by 5 to obtain the mount of nonpreciptive sole digested hemoglobin produced in one minute instead of in 5 minutes; it must be divisted by 5.18 it the molecular weight of tyrosuse of bothan the tyrosuse equivalent as millicquivalents instead of as milligrams. The relation between the number of pepan units in 1 ml of enzyme solution and the colorimeter reading V is thus

$$P \ U = \left[\frac{20}{\sqrt{0.15 + 0.015}} - 0.015 - 0.01\right] \frac{16}{3} \times \frac{1}{5} \times \frac{1}{181} = \frac{0.0104}{\sqrt{1}} - 0.000147$$

¹¹ Northrop J Gen Physiol 16 41 (193°)

¹³ Bubble CO through whipped or blood Centriuge Siphon off the serum and cells. Wash the corpuscles four times with old 9 per cent NaCl. Add an equal volume of water and a sixth of the total volume of tolen? Let stand in the college of the c

stant shaking add 100 ml of the casein solution. Any turbidity must quickly disappear Add different known amounts of pepsin containing solution to the three flasks and make up to 300 ml Keep ut 40° C for one hour Add 20 per cent sodium sulfate solution to the 400-ml mark. Filter and titrate 100 ml of filtrate with 6 I N NaOII, using phenolphthalein as an indicator.

CALLIATION Multiply the litration by 4 Correct for blank run without pepsin and for any acidity of the pepsin solution. The square of the result gives units of pepsin in any pepsin method it is best to determine the amount of unknown required to bring about in a definite time the same amount of digestion as a definite amount of a standard solution of pepsin ²⁴

5 Determination of Tryptic Activity Trypsin is not a gastric enzyme but occurs in the pancreatic juice (see p. 390). In case of regurgitation of intestinal contents through the pylorus, trypsin would be passed into the stomach. The regurgitation is doubtless of frequent occurrence and may even be a normal mechanism by which gustric acidity is regulated (see p. 378). Trypsin is, therefore generally present in the contents of the normal human stomach. However, insamuch as trypsin is distributed by the pepsin-lydrochloric acid of the gastric juice determinations of this enzyme must be carried out immediately after aspirations of the gastric contents particularly where the acidity of the latter is high.

Spencers Method 27 (a) Prepare five reagent tubes, Nos. 1, 2, 3, 4, and 5, more if desired

To Tubes 1 and 2 add 0 5 ml of gastric contents (filter if cloudy)

(b) To Tubes 2, 3, 4, and 5 add 0 5 ml of distilled water

(c) From Tuhe 2 remove 0 5 ml of its mired contents and add to Tube 3
Mix thoroughly and add 0 5 ml from Tube 3 to Tube 4 Repeat for Tube 5

We now have dilutions of gastric contents of 1, ½, ¼, ½, and ½ 6 (d) To each tube add one drop of phenoiphthalein solution (phenoiphthalein 1g, alcohol (95 per cent) 100 ml), then drop by drop add a 2 per cent

lein 1 g, alcohol (95 per cent) 100 ml), then drop by drop add a 2 per cent aodium bicarbonate solution until a light pink color is produced (e) To Tubes 1, 2, 3, and 4 add 0 5 ml of casein solution Tube 5 must re

ceive 1 mi of casem solution, since it contains 1 mi of the diured gastric contents. For the casem solution, dissolve 0.4 g of caseln in 40 ml of 0.1 NaOl1 Add 130 ml of distilled water, then 30 ml of 0.1 NICI. This leaves the solution alkaline to the extent of 10 ml of 0.1 NaOl4, minus about 3 ml neutralized by the casein

(f) Incubate for five hours at 40° C

(g) Precipitate the undigested case in by the dropwise addition of a solution of the following composition glacial acetic acid 1 ml, alcohol (95 per cent) 50 ml, distilled water 50 ml. The tubes in which digestion has been complete remain clear, others become turbid

(h) The tryptic values are expressed in terms of dilution. Thus, complete digestion in Tube 3 (a dilution of 14) shows four times the tryptic power of undiluted gastric julce, taking this dilution as unity, the undiluted juice has a tryptic value of 4.

²⁷ Elaborated by Spencer in the sensor author a laboratory for the specific purpose of determining trypsin in gastrie junce

¹³ Introduce 100 g of pure casem into a 2-liter flask. Add 1000 ml of water shake and let atand some hours. Add 80 ml of N NoH make up to 2000 ml an i warm gradually until clear. Then he al rapidly to 8.5° 00° C to destroy amy proteinases and preserve in a stoppered bottle with a little tolus! Thus solutions keeps for a long time.

[&]quot;Treat 10 g. of a good person preparat in with 100 ml of 10 per cent \aCl Let stand one week at room temperature Filter Add an equal volume of gly cerol an I keep in the ice box The solution keeps indefinitely

- (f) Controls of boiled gastric contents plus casein solution, and of distilled water plus casein solution, treated as above stated, must show no digestion, and become turbid on addition of the precipitating solution
- 6. Detection of Lactic Acid When the audity of the stomach contents is reduced to a low value there may occur considerable fermentation of carbohy drates which have been introduced into the stomach in the ingested food This fermentation yields various organic acids, among which lactic acid is particularly prominent. It is important, therefore, in case of low gastric acidity that the stomach contents be examined for lactic acid.

ETREN-FERRIC CHLORIDE TEST (STRALSS) A satisfactory deduction regarding the presence of lactic acid can be made only by removing the lactic acid from interfering material (e.g. hydrochloric acid protein digestion products etc.) present in the stomach contents Lactic acid may be extracted from the stomach contents by ether. The following technique not only serves to detect lactic acid but also gives an approximate idea as to the amount of the acid present.

Procedure introduce 5 ml of strained stomach contents into a small graduated separatory funnel, add 20 mi of ether, and shake the mixture thoroughly Permit the ether to separate, then allow all the fluid to run out of the separatory funnel except the upper 5 ml of ether To this ether extract add 20 ml of distilled water and 2 drops of a 10 per cent solution of ferric chloride and shake the mixture gently A slight green color is obtained in the presence of 0 05 per cent factic acid whereas 0 1 per cent factic acid yields a very intense yellowish-green color

7 Detection of Occult Blood 13 Benziding Reaction This is one of the most delicate of the reactions for the detection of blood Different benziding preparations vary greatly in their sensitiveness, however. Inasmuch as benziding solutions change readily upon contact with light at is essential that they be kept in a dark place. The test is performed as follows To 3 ml of a saturated solution of benziding in glacial acetic acid: add 2 ml of the solution to be tested and then 1 ml of 3 per cent hydrogen peroxide. A positive test is indicated by a green or blue color.

CONFIDMATORY Test II the mixture contains fat, make neutral or slightly alkaline, and extract by shaking with an equal volume of ether Discard this ether extract Make the residue acid with acetic acid and again extract with ether Pour off ether into a small evaporating dish Put on a hot water bath with flame turned out and evaporate to dryness Add a few drops of water, a drop of benzidine solution, and a drop of 3 per cent hydrogen peroxide A green or blue color indicates blood

Blood is found in gastric contents in conditions associated with erosion of the mucous membrane, ulcer, and curcinoma. In cases of ulcer the blood may be bright red or may be converted to brown "acid hematin" by the excess of acid which is usually present. In carcinoma the blood forms brownish black lumps, the so-called "coffeeground" meteral.

8 Detection of Bile in Stomach Contents If we accept Boldyrell's theory as to the automatic regulation of gastric acidty under normal conditions by the regurgitation of alkaline material from the intestine, then the presence of bile in the gastric puce does not possess the clinical significance it has been accorded. However, if an

[&]quot;These tests may be made upon the strained stomach contents or upon the solid residue
"Glacial acetic acid is preferable, but alcohol acidified with acetic acid may be used

quantitative data are available for seven of these. Although earlier ultra centrifugal measurements indicated a molecular weight of 2000 the values for total sulfur and cystine suggest a figure twice this. The activity of sceretin is destroyed by pepsin and trypsin lience it cannot be given effectively by mouth Secretin is likewise destroyed on incubation with blood scrum presumably because of the presence of the enzyme sccretinase

Harper and Raper first determined that extracts of duodenal mucosa contain two hormonal agencies affecting the external secretory activity of the pancreas These are (1) secretin which stimulates the production of fluid and bicarbonate by the panereatic acini and (2) panereozymin which governs the production of the panereatic enzymes Panereorymin stimulates the output of all panereatic enzymes and when injected repeatedly sustains high enzyme concentrations for prolonged periods Like sceretin pancreozymin is inactivated by blood scrum. Using dogs with a part of the pancreas subcutaneously autotransplanted Wang and Grossman2 have observed that products of protein digestion (peptone and amino acids) are most effective in stimulating release of panereozymin next most effective are soaps (sodium oleate) and to a lesser exter t HCl Carbohydrates are ineffective

It might be supposed that a failure of the pancreatic secretion would be noted in achylia in which condition no gastrie acid is secreted to bring about the liberation of secretin This is not the ease Tatty acids formed in fat digestion and protein split products may cause the passage of secretin into the blood stream. For the same reason bile also has some stimulating action on panereatic secretion

PANCREATIC JUICE

The juice as obtained from a permanent fistula differs greatly in its properties from the juice obtained from a temporary fistula and neither form of fluid possesses the properties of the normal fluid Pancreatic juice collected from a natural fistula has been found to be a colorless clear alkaline fluid (pH 8 or thereabouts) which foams readily. The inorganic salts consist largely of sodium chloride and biearhonate in approximately equal amounts the bicarbonate obtained in part from blood but largely from metaholic processes within the pancreas is responsible for the alkalimity of the juice It is further characterized by containing albumin globulin proteose and peptone nucleoprotein is also present in traces The average daily secretion of pancreatic nuice is 650 ml and its specific gravity is 1 008 The fluid contains 13 per cent of solid matter and the −0 47° C freezing point

The normal panercatic secretion contains a variety of different enzymes Among those which have been well characterized are included the following (1) The peptide-splitting enzymes trypsin chymotryps n and various carboxypolypeptidases (2) a polynucleotidase (or nuclease) acting to split nucleic acids into their component mononucleotides (3)

Harper and Raper J Physiol 102 115 (1913)
Wang and Grossman Am J I hysiol 164 527 (1951)
Hollander and Burnbaum Trans h Y Acad Srs 15 56 (1952)

pancreatic amylase (amylopsin), an amylolytic enzyme, and (4) pancreatic lipase (sterpsin), a fat-splitting enzyme Other enzymes are doubtless present. It will be noted that pancreatic juice contains enzymes capable of acting on all three classes of foodstuffs—proteins, carbohydrates, and fats. It y and co-workers have reported, however, that the concentrations of these enzymes in pancreatic juice can be markedly influenced by diet. Thus, animals receiving a high protein diet secrete a juice especially rich in trypsin, on a high carbohydrate intake the pancreatic amylase concentration is greatly increased, on a high fat-low protein diet, however, the secretion of all enzymes is inhibited. It has been suggested that carbonic anhydrase is importantly related to the intracellular conversion of CO₂ to H₂O₃ and hence to the formation of HCO₃ for secretion.

Proteolytic Enzymes of Pancreatic Juice. The major proteolytic enzymes of pancreatic juice appear to be (1) trypsim, (2) chymotrypsim, and (3) a carboxypolypeptidase, each of which has heen obtained in crystalline form The combined activities of these (and possibly other) enzymes was formerly behaved to he due to a single enzyme which was called trypsin This term is now, however, applied to a single enzyme in the group As a class, these enzymes are similar in that they all catalyze the hydrolytic splitting of the peptide hond They differ from one another in such respects as the type of peptide linkage required for activity, the type of activation from zymogen precursors, as discussed helow, the pH optima, etc

Trypsin and chymotrypsin may be classified as endopeptidases (Bergmann) acting on peptide linkages either in the central portion or the terminal portion of polypeptide chains By the use of synthetic peptide substrates, Bergmann and Fruton found that trypsin acts on peptide linkages containing the carboyl group of either lysine or arginine Chymotrypsin on the other hand was found to act on peptide linkages involving the carboyl group of tyrosine and phenylalanine. Thus the digestive action of trypsin and chymotrypsin on proteins appears to involve the splitting of specific types of peptide linkages in the molecule, the products of the action being, as with pepsin (see p. 364) either low-molecular-weight polypeptides or free amino acids, depending upon the location of the peptide linkages acted upon. There is some evidence of a synergistic action of these various endopeptidases, the action of one may uncover a binkage previously inaccessible to another.

Trypsin has its greatest activity at pH 8 to 9, the optimum pH depending somewhat on the nature of the substrate. It has some action in weakly acid solution in acid solutions it resists temperatures near the boiling point, the denaturation that occurs being rapidly reversible on cooling Trypsin is fairly stable in acidities as high as pH 2, but is digested by pepsin in acid solutions.

The earbovypolypeptidase activity of panereatic juice is doubtless due to a mixture of enzymes, of which only one has been ery stallized (Tig 97) Carbovy polypeptidase is an example of an exopeptidase. It cataly zes the splitting of a peptide linkage involving the amino group of an amino acid whose carbovyl group is free, i.e., not combined in peptide linkage with another amino acid. The action of carbovypolypeptidase is thus

ordinary I wald meal be fed and bile in any considerable quantity be found throughout the entire course of digestion it may indicate pathologically, a stenosis below the level of the common bile duct. The presence of bile is indicated by a yellowish or greenish color of the specimen el anging to a bright green on standing

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16

Pancreatic Digestion

As soon as the food mixture leaves the stomach it comes into intimate contact with the hile and the panereatic juice. Since these fluids are alkaline in reaction and bave high buffering qualities, there can obviously be little further peptic activity after they have become intimately mixed with the clyme and bave neutralized most of the acidity previously imparted to it by the bydrochlorie acid of the gastric juice. The pancreatic juice reaches the intestine through the duct of Wirsung which usually joins with the common bile duet before opening into the intestine near the pylorus.

STIMULATION OF PANCREATIC SECRETION

Secretin and Pancreozymin. Prior to the work of Bayliss and Starling in 1902 it was believed that substances in the intestine stimulated panercatic secretion through local reflexes from the intestine to the pancreas. This view was based on the observation that the introduction of acid into the intestine was followed by secretion. Bayliss and Starling, however, observed that introduction of acid into a jejunal loop which had been denervated as far as possible still induced secretion. This indicated that the exciting agent acted by way of the blood. Injection of acid direetly into the blood did not affect secretion, which must therefore he induced by some substance given off from the mucosa. A substance capable of stimulating secretion by intravenous injection could be extracted from the intestinal mucosa by acid but not by water. Apparently the mucosa is stimulated by the acid of the gastrie chyme to liberate secretin which passes by way of the blood stream to the pancreas eausing it to secrete. This view has been confirmed by many observers, and most clearly by Ivy, Farrel, and Lucth, who found acid applied to transplanted intestinal loops to cause secretion in the transplanted pancreas. The fact that secretin is more readily extractable from the intestinal mucosa by acid than by neutral solvents was interpreted by Bayliss and Starling to mean that the hormone exists in the mucosa as prosecretin. It seems more likely, however, that secretin exists preformed and that the acid chymc serves to release it from its adsorbed state on the colloidal components of the mucosa.

Secretin was the first hormone to have its function clearly established. Tests on the purest secretin preparation so far obtained show it to be a polypeptide, with generally basic properties. Edman and Agren have demonstrated the existence of 15 (or 16) amino acids in secretin, and

confined to the terminal portion of a polypeptide chain. If the free car boxyl group is blocked, as by the formation of an ester, the enzyme becomes mellective Carboxypolypeptides has a pill optimum around 70

ACTIVATION OF TRY FSIN AND CHANGER FSIN Panereatic juice obtained from a fistula commonly shows no action on printing On entrance into the intestine the juice immediately becomes active Apparently tryp in and chymotrypsin as found in the panereas and as secreted by that gland exist not as such but as proenzy mes called trypsingen and chymotrypsingen. These proenzy mes have been obtained by Northrop and Kunitz

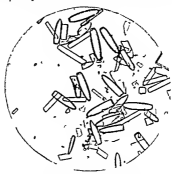


FIG 97 CRYSTAILINE CARSOXYPOLYPEPTIDASE, PREPARED BY THE METHOD OF ANSON Courtesy Dr A A Plenti

in crystalline form and show protein characteristics. Trypsin appears to be normally activated by the substance enterol inace secreted by the intestinal nuccoa. Kunitz has clearly demonstrated that enterokinase acts on trypsinogen enzymatically. Furthermore it should be noted that erystalline trypsinogen (Fig. 98) can be activated by traces of trypsin the activation mereasing in rapidity as more active trypsin is formed. In less pure trypsin preparations activation appears to be slowed up by the presence of an inhibitor which has been obtained in crystalline form and appears to be a polypeptide. Strong salt solutions may also activate trypsin.

Chymotrypsinogen is apparently activated by active trypsin Enterokinase thus does not activate chymotrypsinogen directly but indirectly through its action on trypsinogen

Pancreatic Amylase. This is an amylolytic enzyme or more probably a mixture of enzymes, which possesses somewhat greater digestive power than salivary amylase. As its name implies its activity is confined to the

starches and similar compounds, and the products of its amylolytic action are destrins and, ultimately, maltose.

It is probable that the saliva as a digestive fluid is not absolutely essential. The salivary amylase is destroyed by the hydrochloric acid of the gastric juice and is therefore inactive when the chyme reaches the intestine. Should undigested starch be present at this point, however, it would be quickly transformed by the active pancreatic amylase This enzyme is not present in the pancreatic juice of infants during the first few weeks of life, indicating that a starch diet is not normal for this period.

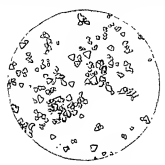


Fig 98 Trypsingen Crystals, Courtesy, Dr John H Northrop,

Meyer and co-workers have reported that pancreatic amylase appears to be identical with salivary amylase. They prepared crystalline amylases (see Fig. 74, p. 308) from hog pancreas, human pancreas, and human saliva, and compared them to ascertain whether enzymes of similar action from different glands of one species are more alike than corresponding enzymes from the same gland of different species. In all respects the two human amylase preparations were indistinguishable, and different from that of the hog pancreas.

Pancreatic amylase possesses protein properties. It requires 1011 such as Cl⁻ or Br⁻ for normal activity, and its optimal pH ranges from 6.3 to 7.2 in the presence of different neutral salts.

It has been shown that pancreatic amylase will digest raw starch. The raw starch of corn and wheat may be completely digested and absorbed by normal adults in amounts of more than 100 g. per day, whereas the raw potato starch is about 80 per cent available.

Pancreatic Lipase. This is a fat-splitting enzyme. It has the power of splitting the neutral fats of the food by hydrolysis, into fatty acid and glycerol. A typical reaction would be as follows:

⁴ Meyer, Fischer, Bernfeld, and Duckert: Arch. Brochem , 18, 203 (1948).

$\begin{array}{c} C_2H_3(\mathrm{OOCC_{13}H_{21}})_2 + 3H_2O \rightarrow 3(C_{14}H_{12}\mathrm{COOII}) + C_1H_3(\mathrm{OII})_1 \\ \text{Palmitin} \\ \text{Palmitic acid} \\ \end{array}$

Pancreatic lipase is very unstable and is easily rendered inert by the action of acid. For this reason it is not possible to prepare an extract having a satisfactory fat-splitting power from a pancreas which has been removed from the organism for a sufficiently long time to have become acid in reaction.

Fodors has reported that, in addition to lipase, hog pancreas contains a monovalent alcohol esterase. Although the different activities have not as yet been localized in separate protein fractions, the trigly ceride-splitting activity is generally more resistant to heat, alkali, and try psin than is the esterase activity.

Pancreatic lipase is undoubtedly the most important fat-splitting enzyme in the digestive tract. In the absence of paniereatic lipase, for example, when the pancreatic duct is obstructed by disease, the fat of the diet appears in the undigested form in the feces, this condition is known as steatorrhea The enzyme is water-soluble and presumably a protein, although sufficiently pure preparations to prove this have not as yet been obtained, nor has the enzyme been crystallized Its optimum pH is about 7 The action of the enzyme on fats is obviously dependent in large measure upon the surface of fat available, thus the more highly emulsified the fat 19, the more surface is exposed to the aqueous phase containing the enzyme, and the more rapid is lipase action Emulsification of fats in the digestivo tract is facilitated by the presence of bile, because of the lowering of surface tension brought about by the bile salts (see Chapter 18) The bile also aids in fat digestion by facilitating removal of the end products of lipase action It is believed by some that the bile actually activates pancreatic lipase, it appears more probable that the acceleration in hoase activity in the presence of bilo is due to physiocochemical action by the bile salts in facilitating closer contact between the water-soluble lipase and the fat globule

PREPARATION OF AN ARTIFICIAL PANCREATIC JUICE

After removing the fat from the pancreas of a pig or sheep, finely divide the organ by means of scissors and grind it in a mortar if convenient, the use of an ordinary meat chopper is a very satisfactory means of preparing the pancreas

When finely divided as above, the pancreas should be placed in a 500 ml flask, about 150 ml of 30 per cent alcohol added, and the flask and contents shaken frequently for 21 hours (What is the reaction of this alcoholic extract at the end of this period, and why?) Strain the alcoholic extract through cheescoich, filter, nearly neutralize with potassium hydroxide solution, and then exactly neutralize it to litmus with 0.5 per cent sodium carbonate

PRODUCTS OF PANCREATIC DIGESTION OF PROTEIN

Into a 250 mi flask introduce 20 g of casein, 10 mi of the artificial pan creatic juice prepared as described above, and 100 ml of 1 per cent sodium car bonate Allow to digest at 40° C for 8 to 10 days with the addition of a few milliliters each of chloroform and toluene, the flask being stoppered with

Fodor Arch Buchem 26 307 (1950)

cotton. As the chloroform and toluene evaporate they must be renewed. Heat the mixture to boiling, and at the boiling point add accetic acid drop by drop until the mixture is acid in reaction. Cool and filter.

To 5 ml. of the filtrate add bromme water drop by drop. Note the development of a pink color which disappears in the presence of an excess of the

reagent. This reaction indicates the presence of tryptophan.

To another 5-mi. portion of the filtrate add 10 drops of concentrated sulfuric acid and 10 ml. of a 10 per cent solution of mercuric sulfate in 5 per cent sulfuric acid. After mixing and allowing to stand for a few minutes, filter of the yellow precipitate which forms. This is an impure mercury compound of tryptophan. Filter off the precipitate, reserving the filtrate, and wash the precipitate on the filter paper thoroughly with several small portions of water.

To small portions of the precipitate apply the Hopkios-Coie, xanthoproteic, and Milion tests. Tryptophao gives a positive reaction with the first two of these tests and is responsible for the Hopkins-Cole reaction as applied to protein.

Test portions of the filtrate from the mercuric precipitate hy the Hopkins-Cole, xanthoproteic, and Millon reactions Tyrosine responds to the latter

T AL

To the remainder of the original filtrate add a few drops of ammonia' (enough to make it siightly alkaline) and evaporate to a volume of 10 to 20 mil, using at first a free fame and completing the evaporation on a water bath. Transfer to a beaker and allow to stand for 1 or 2 days. Examine microscopically the crystals which separate out. Tyrosine crystallizes in sheaves of needles (see Fig. 44). Leucine forms small rosettes. Apply Mörner's reaction for tyrosine (see p. 138).

GENERAL EXPERIMENTS ON PANCREATIC DIGESTION

EXPERIMENTS ON TRYPSIN⁸

- 1. The Most Fovorable Reaction for Tryptic Digestion. Prepare five tubes as follows:
 - a. 3 ml. of neutral pancreatic extract + 3 ml of water, pll 7.
- b. 3 mi. of neutral pancreatic extract + 3 ml. of water + 1 drop of phenoiphthalein solution + 0 5 per cent Na₂CO₂ to first faint piok color. pH 8.3.
- c. Same as (b) but add Na.CO, until the pink color no longer deepens. pH 10.
- d. 3 mi. of neutral pancreatic extract + 3 ml. of 2 per cent boric acid solution. oil 5
 - e. 3 mi. of neutral pancreatic extract + 3 mi. of 0 6 per cent IICi. pli 3.
- e. 3 mi. of neutral pancreatic extract + 3 mi. of 0 6 per cent HGL pli 3. Add a small plece of fibrin' to the contents of each tube and keep them at 40° C., noting the progress of digestion. In what reactions does trypsin act

It has been claimed that a similar yellow precipitate forms in the presence of tyrosine

c) stime and polypeptides

7 If the solution is alkaline in reaction owing to the presence of fixed alkali while it is being concentrated the amino acids will be broken down and ammonia will be liberated. Ammo-

nia in slight excess does not cause such decomposition

1 I or these experiments as well as for those on the other pencreatic enzymes commercial

preparations of it; youn and pancreatin may be employed

1 Congo red fibrus (see Appendax) may be used in this and the following tests on try ptic
digestion: If Congo red fibrus (see used the experiments should be carried out at room temperature Also in Exp. (b) and (c) phenolophthalen should not be added, but the proper
amount of alkali as determined in a separate test. Buffer solutions of suitable pII may also
be used in these tests instead of the acids and alkalies suggested.

and what is the optimum pli? liow do the indications of the digestion of fibrin by trypsin differ from the indications of the digestion of fibrin by pep sin? Is the same degree of swelling of the protein noted?

- 2 The Most Favorable Temperature (For this and the following series of experiments under tryptic digestion use the neutral extract plus an equal volume of 0 5 per cent sodium carbonate ; In each of four tubes place 5 ml of alkaline pancreatic extract Immerse one tuhe in Ice water, keep a second at room temperature, and place a third in the incubator or water hath at 40° C Boil the contents of the fourth for a few moments, then cool and also keep it at 40° C into each tube introduce a small plece of fibrin and note the progress of digestion. In which tube does the most rapid digestion occur? What is the reason?
 - 3 Demonstration of the Action of Enterokinase on Trypsin

a Preparation of Enteroarnast Grind 5g of fresh duodenal mucosa of the hog with a little sand Gradually add 50 ml of water during the grinding process Strain through cheesecloth

A better preparation is made hy drying the mucosa From the upper three feet of the intestine of the hog scrape off the mucous membrane with a knife or, hetter, a glass plate Shake with three volumes of acetone Let stand two hours Fliter Wash the residue again with the same amount of acetone, then with a mixture of acetone and ether, and finally twice with ether Dry in air and pulverize A 1 50 extract of this powder in water may be used The powder keeps indefinitely The enterokinase may be further purified

b Preparation of Kinase-free Trypen Solution Immediately after killing the animal grind hog pancreas in a meat chopper and dry with acetone and ether as in preparation of enterokinase (see above) Glycerol extracts of this dried pancreas (1 10) may be used Kinase free trypsin may also be further purified

- e DEMONSTRATION OF ACTION OF ENTERORINASE Prepare five tubes as follows
- (1) 2 ml of pancreas extract + 5 ml of water
- (2) 2 ml of pancreas extract + i mi of duodenal extract + 4 ml of water
- (3) 1 ml of duodenal extract + 6 mi of water
- (4) 2 ml of pancreas extract + 1 mi of duodenal extract + 4 ml of water (5) 2 ml of pancreas extract + 1 ml duodenal extract (boiled) + 4 ml ol water

Boil contents of Tube 4 for 5 minutes and cool to 40° C Keep all tubes at 40° for 20 minutes for activation Add 1 ml of 0 5 per cent sodium carbonate to each tube and the same quantity (about the size of a pea) of fresh fibrin Shake the tubes and place at 40° C Observe frequently during the course of an hour Tube 2 should show the most rapid digestion Why?

4 Quantitative Determination of Traptic Activity 19 See Spencer & Method p 390

EXPERIMENTS ON PANCREATIC AMYLASE

1 Demonstration of Presence of Amylase in the Pancreas Into a test tube introduce 5 ml of starch paste and 2 ml of pancreatic extract Shake and put in a water bath at 40° C for 30 minutes Divide into two parts Test one with lodine for undigested starch and the other hy Benedict s test for reduc

A number of proteclytic enzymes may be quantitatively determined by means of the fit in plate method. This procedures is by the convenient and sens tive. Astrop and Alkjaersis. Arch. Biochem. Bioph. 37, 93 (19.2)

ing sugar. The reducing sugar formed by pancreatic amylase is maitose as is the case with salivary amylase.

2. Quantitative Determination of Amylolytic Activity: Method of Willstatter, Waldschmidt-Leitz, and Hesse. In this method the reducing sugar formed is determined by hypolodite titration.

Into a 50-ml, cylinder (with a ground-in stopper) introduce 25 ml, of a freshly prepared I per cent solution of soluble starch (see the Appendix). 10 ml, of buffer solution pH 6.8 (5 1 ml, of 0 2 M KH, PO, + 4.9 ml, of 0.2 M Na. HPO.), and 1 ml, of 0.2 N NaCl. Mix and bring to a temperature of 37° C. Add the enzyme solution to be tested. Mix and return at once to the bath Keep at 37° C. for exactly ten minutes Then add 2 ml. of N HCl to stop the action. Wash the contents of the cylinder with a little water into an Erlenmeyer flask. Add 0.6 ml. of 0.1 N todine solution for each me, of maltose expected. Then odd drop by drop with shaking 0.1 N NaOH sufficient to neutralize the added HCl and to change the primary phosphate of the huffer to secondary form (for both of which 30 ml is required) and a further amount 1.5 times the volume of the jodine solution used. Let stand 15 minutes. acidify with dilute H1SO4, and titrate the excess iodine with 0.1 N thiosulfate. The iodine taken up by the starch and the enzyme solution is found in a control determination. One ml. of 0 1 N I is equivalent to 17.15 mg. of Gi-H-On or maltose.

CALCULATION. The equation for a monomolecular reaction is

$$k = \frac{1}{t} \log \frac{a}{a - x}$$

Apply this formula to the determination above a is the amount of substrate (oot the full 0.25 g of starch but 75 per cent of this or 0.1875, as representing the practical limits of saccharification of the starch), t is the time (10 mioutes). Assume the iodine required to be 2.29 ml and the blank determination 0.53 ml. Then the iodino taken up by the maltiose w ill be 2.29 - 0.53 = 1.76 ml of 0.1 N I, equivalent to 0.0302 g of maltiose, and

$$L = \frac{1}{10} \log \frac{0.1875}{0.1875 - 0.0302} = 0.0076$$

This reaction constant expresses also the number of units of amylase in the amount of amylase preparation used. The unit of amylase is 100 times the amount required under the given conditions to give a constant of 0.01. This is approximately the amount in 2 cg. of dired punciess. The amyloly to strength of a preparation may be expressed in units per cg. For plant amylase a buffer of pH 5 should be used. For accurate work the amount of enzyme used should be such as to give a constant of 0.01.—0.03 or hydrolysis of not over 40 per cent of the substrate.

EXPERIMENTS ON PANCREATIC LIPASEIT

- Influence of Bile on Action of Lapase Prepare five test tubes as follows

 0 5 ml
 of olive oil + 5 ml
 of neutral pancreatic extract + 4 5 ml
 of olive oil + 5 ml
 - b. 0.5 ml, of olive oil + 95 ml of water.
 - c. 9.5 ml. of olive oil + 85 ml of water + 1 ml of blie.
- d. 0.5 ml of olive oil + 5 ml. of neutral poncreatic extract + 3 5 ml. of water + 1 ml of bile
 - e. 5 ml, of neutral pancreatic extract + 4 ml, of water + 1 ml, of bile.

¹² A vegetable lipase preparation may be made as described on p. 331

Shake the tubes thoroughly, add a drop of toluene to each, and place them in an incubator or water hath at 40° C. for 24 hours. At the end of this period add a drop of phenoiphthalein to each tube and titrate with 0 65 N NaOii to a permanent pink color. Shake the tube during the titration Record the amount of 0.05 N alkali necessary to neutralize the contents of each tube. Which tube required the most? Wby?

- 2. Litmus-miik Test. Into each of two test tubes introduce 10 ml. of milk and a small amount of litmus solution.¹² To the contents of one tube add 3 ml. of neutral pancreative extract: and to the contents of the other tube add 3 ml. of water or of boiled neutral pancreatic extract. Keep the tubes at 40° C. and note any changes which may occur. What is the result and how do you explain it?
- 3. Copper Soap Test for Lipase. Prepare a 2:100 agar-agar solution, mix with an equal volume of 5:100 starch paste, incorporate in this mass about 1/40 of its volume of the neutral fat desired (butter, lard, etc.), heat with constant agitation until a homogeneous emulsion is produced, pour into a Petri dish, and cool rapidly. With a fine pipet distribute on the surface of the solidified mass small drops of the liquid to be tested, keep 1 hour at 38° C, pour a saturated aqueous CuSO, solution over the surface, allow to stand 19 minutes, and rinse with H₂O. The presence of lipase is shown by the appearace of beautiful bluish-greeo spots. These are copper soap. The addition of the starch, which is not indispensable, produces a rather white opaque back-ground against which the spots appear very distinct.
 - 4. Ethyl Butyrate Test. Into each of two test tubes introduce 4 ml. of water, 2 ml of ethyl butyrate, Gill-COO Gill, and a small amount of litmus powder. To the cooteots of ooe tube add 4 ml. of neutral pancreatic extract and to the contents of the other tube add 4 ml. of water or of holled neutral pancreatic extract. Keep the tubes at 40° C. and observe any change which may occur. What is the result and how do you explain it? Write the equation for the reaction which has taken place.
 - Quantitative Estimation of Lipase Method of Willstätter, Waldschmidt-Leltz, and Memmen. A constant degree of activation is obtained by the addition of albumin and calcium salts. An adsorption compound fat-Ca oleatealbumio-lipase very favorable to lipase action is believed to be formed.

into a wide-mouth 30-ml. flask with a ground glass stopper, introduce the enzyme preparation made up to 10 ml. with water, then 2.5 g. of olive oil and 2 mi of buffer (0.66 ml. of N NII.011 + 1.34 ml. of 1 N NII.021) and 0.5 ml of 2 per cent CaCl₃. Shake a little and add 0.5 ml of 3 per cent albumin solution-shake by hand uoliormly aod strongly for 3 minutes to get a stable characteristic emuision. Put in a thermostat at 30° C. for 57 minutes (total time of lipase action 00 minutes). Wash into an Erlemmeyer flask with 96 per cent alcohol to make a volume of about 125 ml. Add 20 ml of ether and mix. This stops lipase action. Add 12 drops of 1 per cent solution of thymolphthalen and stirate with 0.1 N alcoholic KOII to a distinct blue color. From the read nig subtract the titration value of the huffer (13.4 ml of 0 t N KOII) and of the enzyme solution A lipase unit is the amount required under the specified

¹³ Litmus-milk powder may be used if desired. To prepare it add 1 part of powdered litmus to 50 parts of dired milk powder. For use in testing, 1 part of powdered litmus-milk may be added to 9 parts of water.

is Commercial pancreatin may be used in this test if desired

conditions to split 24 per cent of 2 5 g of ollve oll of a saponification value of 185 (meaning that 1 g, of the ollve oll completely hydrolyzed requires for neutralization 185 mg, of KOH)

CALCULATION One ml of 0.1 N KOH = 5.61 mg of KOH If 12.5 ml of 0.1 N KOH (corrected for control) were required $12.5 \times 5.61 \approx 70.6$ mg of KOH Then 70.6×100 = 15.1 per cent split If the splitting is more than 24 or less than 10

 $\overline{185 \times 25} = 16$) per cent spitt if the spitting is more than 24 or less than 10 per cent repeat. Ten per cent decomposition by panercatic lipase corresponds to a lipase value of 0.028, and 24 per cent to 10 Add to 0.28 an amount of 0.044 for each per cent of decomposition above 10 and up to 24. If using other than panercatic lipase or if other conditions are altered a special curve of variations of percentage hydrolysis with different amounts of enzyme must be plotted from experimental data obtained using the enzyme in question. Concentration of lipase may be expressed as units per ego of substance

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17

Intestinal Digestion

The digestive processes carried out by the enzymes of the paneratic juice occur within the lumen of the intestine. It is probable that peptic digestion also continues in the upper part of the duodenum until the acid chyme has been neutralized by the paneratic and intestinal juices. However, we shall consider in this chapter only those digestive processes which are effected by enzymes secreted in the intestinal juice (succus enterious) and those present in the intestinal microsa.

The intestinal juice is most abundant in the duodenum and jejunum and is of two types (a) a sceretion poor in enzy me content but rich in mucin which is produced periodically at about two-hour intervals, and (b) a digestive juice which is secreted in response to a meal, probably through mechanical stimulation of the intrinse nerves of the intestine. A hormone entercornin, liherated by the mucosa in response to a meal,

may be concerned in stimulation of the intestinal glands

The reaction of the intestinal contents is influenced by the state of digestion and the relative volumes of acid chyme, pancreatic and in testinal junces, and hile present in any given segment of intestine The reaction of the small intestine contents is frequently acid, and the pH is usually considerably lower than that of the intestinal junce Intestinal secretions collected at intervals from the joinum through the colon contain about the same concentrations of cations, but progressively higher concentrations of bearbonate and lower concentrations of chloride Thus the intestinal secretions are more alkabne in the lower reaches of the intestinal secretions are more alkabne in the lower reaches of the

The enzymes of the intestine are of great importance, since their action, in some cases, supplements that of other digestive enzymes. They also serve to complete the digestive process by cataly zing the further hydrolysis of products of other digestive reactions. The enzymes include (1) peptidases (aminopolypeptidases and dipeptidases), (2) carbohydrases (cuerase, maltase, and lactase), (3) enzymes acting on nucleic acid-including a nonspecific phosphatase, (4) leathinase Some of these enzymes are not actually present in the intestinal juice, or occur there in only small amounts Examples of such enzymes are lactase nucleotidase and nucleoidase. These enzymes evidently evert their action at the surface of the muco al cells or intracellularly. These latter sites of action may also be important in the case of other meetinal enzymes.

Peptidases Annopolypeptidases of the ammopolypeptidases of the intestinal mucosa act on the products of peptic and tryptic digestion to

catalyze the splitting of the peptide linkage adjacent to the end of polypeptide chains hearing the free amino group. The products are a free amino acid and a polypeptide of lower molecular weight. These enzymes supplement the effect of carbox polypeptidases of pancreatic juice which have their site of action at the carboxyl end of the polypeptide chain. Complete hydrolysis of polypeptides may be effected by aminopolypeptidases, since they slowly hydrolyze synthetic dipeptides, but this result is better accomplished by the dipeptidases. Some structural specificity is undoubtedly myolved in aminopolypeptidase action, thus leucylpeptidase in bog intestinal juice is especially active in hydrolyzing peptides containing leucine as a terminal amino acid.

DIPERTINASES The dipertidases have relatively little action on

polypeptides but split dipeptides rapidly

The combined activity of aminopolypeptidases and dipeptidases was once attributed to a single enzy me called erepsin. This name is now used only as descriptive of the over ill results of these enzymes. Ereptic enzymes are found not only in intestinal microsa but in other plant and animal tissues as well. None of these enzymes has been crystallized. Since the ereptic enzymes (and carboxypolypeptidase) act only on peptide linkages adjacent to the end of a polypeptide or peptide chain, they are referred to as exopeptidases to distinguish them from pepsin and the tryptic enzymes which act on centrally located peptide honds. Through the combined action of all these enzymes, protein is finally by drolyzed to individual amino acids.

Carbohydrases. Sucrase The three carbohydrases sucrase, maltase, and lactase are also important enzymes of the intestinal mucos. The sucrase acts upon sucrose and inverts it with the formation of glucose and fructose Sucrases may also be obtained from several vegetable sources. For investigational purposes sucrase is ordinarily obtained from yeast (see p. 331). Intestinal sucrase has an optimum pH of 5 to 7, yeast invertase acts best at pH 45

Luctase This enzyme splits lactose with the consequent formation of glucose and galactose Freshly prepared suspensions of intestinal mucosul tissue are much more active than water extracts, or the intestinal juice, indicating that the activity is intimately associated with the mucosal cells. The optimum pH for intestinal lactase is about 54 to 60.

MALTASE This enzyme possesses the power of splitting multose, the end product of the digestion of starch by amylase, into glucose Maltase is

best prepared from yeast Its optimum pH is 67 to 72

Nucleases A polynucleotudase in intestinal juice results in the depolymentation and break up of nucleic and to nucleotudes. The nucleotudes are then by droly zed by a nucleotudase (a phosphitase) to give purine and pyrimidine nucleosides and phosphoric and A nucleosidase decomposes the purine nucleosides to the purine base and riboso (or deoxyribose). The pyrimidine nucleosides are absorbed unchanged but are decomposed by enzymes found in other organs. The intestinal phosphoriase responsible for the hydroly sis of the phosphoric and ester linkage of nucleotides is onspecific in its action, since it attacks other phosphoric and ester substrates such as giverophosphate and heose phosphates. A part of the

tryptic enzymes

alkaline phosphatase activity of normal plasma is due to the intestinal phosphatase which has entered the blood

Lecithinases Enzymes which hydrolyze lecithins to various products depending upon the experimental conditions and time of action are

depending upon the experimental conditions and time of action are present in intestinal mucosa Enterokinase This substance is discussed in Chapter 16 Though present in intestinal juice it is not a digestive enzyme but activates the

GENERAL EXPERIMENTS ON INTESTINAL DIGESTION

Preparation of Intestinal Extract Wash a piece of hog intestine about 18 inches long Run it through a meat chopper or, better, scrape off the mucosa with a knife or piece of glass Rub in a mortar with sand Add 5 volumes of water and a little chloroform and let stand at room temperature for 24 hours if necessary to expedite laboratory work the extract may be used after two hours Strain through cloth or loose cotton This extract may be used for the general experiments on phosphatase ereptic enzymes, and sucrase The presence of some mucosal cells in suspension increases the enzyme activity

EXPERIMENT ON INTESTINAL PHOSPHATASE

Demonstration of Actson of Intestinal Phosphatase on Nucleic Acid and on Sodium Glycerophosphate Prepare a 2 per cent solution of yeast nucleic acid with the aid of just sufficient diluxe haOll to make the resulting solution pink to phenolphthalein (pil 8 6) Then prepare a 2 per cent solution of sodium glycerophosphate and make just pink to phenolphthalein To each of four test tunes add 10 ml of the Intestinal extract prepared as above Boll two—Tubes 2 and 4—for one to two minutes To Tubes 1 and 2 then add 10 ml of the 2 per cent nucleic acid solution and to tubes 3 and 4,5 ml of the glycerophosphate solution Add 2 to 3 ml each of toluene and chloroform to each mixture Keep at 38° C for 24 hours

Heat the tubes to boiling in a water bath to coagulate protein Add 5 ml of 5 per cent HCl and allow to stand for one hour This precipitates any un changed nucleic acid Filter and take allquors of the filtrate (about 10 ml) Precipitate the phosphate from each mixture by adding 5 ml of magnesia of magnesium ammonium phosphate should be found in Tubes 1 and 3 indicating that the phosphotic acid of the nucleic acid and of the glycero The control should show only a slight precipitate.

EXPERIMENTS ON PEPTIDASES

1 Demonstration of Peptidase Activity To about 5 mi of a I per cent solution of peptone in a test tube add about 10 mi of the intestinal extract pre pared as described above Prepare a second tube containing a like amount of peptone solution, but boil the intestinal extract before introducing it Place the two tubes at 38° C for two to three days At the end of that period best the contents of each tube to boiling filter and try the bluret test on each filtrate in making these tests care should be taken to use like amounts of distrate londle, and copper sulfate in each test in order that the distract of the contents of the tube which contained the boiled extract should show a deep pink color with the bluret test owing to the peptone still present On the other hand, the bluret

test upon the contents of the tube containing the unboiled extract should be negative or exhibit, at the most, a faint pink or bine color, signifying that the peptone, through the influence of the intestinal extract, has been transformed, in great part at least, into simple peptides and amino acids which do not respond to the bluret test. To other portions of the filtrates add a few drops of hromine water. A violet color indicates free tryptophan and hence that amino acids have been liberated. Proteinase in the extract changes the peptone to polypeptides. The latter are broken down by aminopolypeptidase to simple peptides which under the action of dipeptidase yield amino acids. The action of several enzymes is thus involved.

- 2. Demonstration of Dipeptidase Using Glycyltryptophan. Introduce 5 ml. of glycyltryptophan solution into each of two tubes. To one add 5 ml. of intestinal extract, to the other 5 ml. of boiled extract. Let stand over night. Add a few drops of bromine water to each. A vloiet color indicates free tryptophan and hence the presence of dipeptidase.
- 3. Quantitative Determination of Dipeptidase. To 25 ml. of 0.05 N glycyl-glycine add NaOll to pH 8. Warm to 37° C. Add enzyme solution. Carry out a formol thration on an aliquot at once (control) and repeat at intervals Choose a determination representing less than 30 per cent digestion. Calculate $k \ln i$: $k = (1/t) \cdot \log a/(a-x)$. Activity equals k/g enzyme preparation,

EXPERIMENTS ON CARBOHYDRASES

1. Demonstration of Sucrase. To about 5 ml. of a 1 per cent solution of sucrose, in a test tube, add about 1 ml of intestinal extract, prepared as described above, and a few drops of chloroform. Prepare a control tube in which the Intestinal extract is bouled before being added to the sugar solution. Let stand for 24 hours. Heat the mixture to boiling to coagulate the protein material, filter, and test the filtrate hy Benedict's test (see p. 66). The tube containing the holied extract should give no response to Benedict's test, whereas the tube containing the unboiled extract should give no response to Benedict's solution. This reduction is due to the formation of glucose and fructose from the sucrose through the action of the enzy me sucrase which is present in the intestinal epithelium.

For the preparation of vegetable sucrase see p 331.

 Demonstration of Lactase and Maliase. Repeat the above experiment but use 1 per cent solutions of lactose and maltose. Is there any evidence of lactase and maltase activity?

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18

Bile and Liver Function

Secretion of Bile The bile is secreted continuously by the liver and passes into the intestine through the common bile duct which opens near the pylorus. This process is continuous even during prolonged fasting provided there is no obstruction The ingestion of food, however, increases the secretion Meat is effective in this respect fat less so, and starch and sugar appear to be without effect

Introduction of acid into the duodenum stimulates bile formation sug gesting that secretin is responsible However, injection of secretin prepa rations though leading to a copious secretion of pancreatic juice in one to three minutes produces a much slighter secretion of bile and then only nfter seven to nine minutes Ligation of the pancreatic vein provents this nction on the liver, hence it is suggested that some metabolic product of the pancreas is responsible

Bile salts absorbed from the intestine have a marked power to stimulate hile formation. The passage of bile directly from the liver or through the emptying of the gallhladder serves to induce a further secretion Deliydro-

cholic acid is an especially effective cholagogue

Mechanism of Emptying of the Gallbladder. There appear to be at least two mechanisms active in gallbladder emptying. One of these in volves the contraction of the gullbladder and the other the tone of the sphineter of Oddi at the entrance of the common bile duct into the inter tine Cream or egg 3 olk causes an emptying of the gallbladder apparently by inducing an active contraction of this organ, probably accompanied by o relaxation of the sphineter at the same time. The active agent is the free fatty and liberated on digestion of these foods. The organic ands of fruits and the gostne and have the same effect. In cases of acute cholecystitis fats acid fruits and meets which stimulate acid secretion in the stomach should therefore be reduced in the diet in favor of cereal foods The contraction is opparently brought about through liberation from the intestinal mucosa of a hormone cholecystokinin whose chemical nature is not yet determined Mognesium sulfate promotes evacuation by causing a dilation of the sphineter

Functions of the Bile We may look upon the bile as an excretion as well as a secretion In the fulfillment of its exerctory function it passed such substances as kerthin metallic compounds chokesterol and the decomposition products of hemoglobin into the intestine and in this way aids in removing them from the organism. As a secretion, the bile as 150 materially 11 the digestion and absorption of fats from the intestine by

its emulsifying action on the fats of the diet and by facilitating the absorption of the fatty acids formed by the action of the pancreatic juice. A decreased appetite for fats has been shown in rats after the ligation of the common bile duct. A further important function of the bile is to aid in the absorption of vitamin K. Symptoms of vitamin K deficiency frequently accompany the absence of an active secretion of bile into the intestine.

Composition of Bile. The bile is a ropy, viscid fluid which is alkaline in reaction (pH 78) when it issues from the liver, and ordinarily possesses a decidedly bitter taste. It varies in color in the different animals. the principal variation being vellow, brown, and green. Fresh human bile from the living organism ordinarily has a vellow brown or golden vellow color Post mortem bile is variable in color It is very difficult to determine accurately the amount of normal hile secreted during any given period For an adult man it has been variously estimated at from 500 ml to 1100 ml for 24 hours. The specific gravity of the hile varies between 1 010 and 1 040, and the freezing point is about -0 56° C. As secreted by the liver, the bile is a clear, limpid fluid which contains a relatively low content of solid matter. This secretion has a specific gravity of approvimately 1 010 After it reaches the gallbladder however, it becomes mixed with mucous material from the walls of the gallbladder and this process coupled with the continuous absorption of water and certain other com ponents from the bile has a tendency to concentrate the secretion Therefore the bile as we find it in the gallbladder ordinarily possesses a higher specific gravity than that of the freshly secreted fluid. The specific gravity under these conditions may run as high as 1 040 There is a decrease in inorganic salts owing to absorption, while the concentration increases the content of organic substances Even though it is concentrated in the gallbladder, the bile remains practically isotonic with the blood because the increased content of high-molecular weight bile salt ions is accompanied by a decrease in chloride and bicarbonate ions. The pH of bladder bile may fall as low as 6, as compared to the definitely alkaline reaction of fistula hile

SELECTED ANALYSES ILLUSTRATING THE COMPOSITION OF HUMAN BILE (PARTS PER 1 000)

Constituent	Festula Bile	Blad ler Bile
Water	9,60	S60 0
Solida	210	140 0
Bile acids	57	53.7
Muem and pigments	8.0	11 1
Total lipides	2 9	18 8
Fatty acids	0.8	8 5
Neutral fat	0.8	15
Cholesterol	0.8	57
Phosphatides	0.5	2 2
Inorganic matter	7.4	8.5

The principal organic constituents of the fale are the salts of the bile acids, ble pigments, neutral fits, lecithm, phosphatides, nucleoprotein, mucin, and cholesterol horganic constituents include chiefly sodium chloride, and bicarbonate, some potassium, calcium, magnesium, and sulfate and a trace of phosphate. The metals iron, copper, and zine are ilso frequently present in detectable amounts. The bicarbonate content of hepatic hile is higher than that of serum, the chloride content of biles lower than that of serum, the chloride ion being replaced largely by the organic bile salt ions.

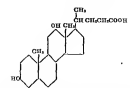
The quantitative composition of bile varies according to the source of the bile, i.e., whether the bile for analysis is obtained from the gallbladder or by means of a fistula before it reaches the gallbladder. The difference in the composition of these two types of bile is shown in the table of

selected analyses on p 409

Bile Acids. The bile scids are elaborated exclusively as far as is known by the hepatic cells of vertebrates. They may be divided into two groups (1) the glycochole acid group and (2) the tainceholic acid group. In human bile glycocholic acids predominate, while taurocholic acids are more abundant in the bile of carnivora. The glycocholic acids are combinations through peptide linkage of bile acids and glycine, NII, CH-COOH. The taurocholic acids are similar combinations of bile acids with taurine, NII, CH; CH; SO; III. Taurine is evidently derived in the body from cysteine.

There are several varieties of bile acids, and therefore there are several forms of glycocholic and of taurocholic acids, depending upon the nature of the bile seid entering into the combination. The principal bile acids are (1) choic acid, C2.H4.004 with three hydroxyl groups, (2) deoxycholic acid C14H40O4 with two hydroxyl groups, (3) anthropodeoxycholic acid and (4) hyodeoxycholie acid which are isomeric with deoxycholic acid differing only in the position of one hydroxyl group, and (5) lithocholic acid C24H40O2 which has a single hydroxyl group Human bile contains the first three of these in the proportion of about three parts of cbolic acid to one part of deoxycholic and some anthropodeoxycholic acid Ox bile contains about 6 parts per 100 of choic acid and about one-eighth as much deoxycholic acid Hyodeoxycholic acid is found in hog bile and chenodeoxycholic acid (identical with anthropodeoxycholic acid) in the bile of the goose and chicken The choice acids are closely related in structure to cholesterol and are probably formed in the body from cholesterol or one of its immediate metabolic precursors. See the structural formulas for cholic and deoxycholic acids

Choile acid (3,7,12 Trihydroxycholanic acid)



Deoxycholic acid (3,12-Dihydroxycholanic acid)

The bile acids are present in the bile largely as sodium salts. The sodium glycocholate and taurocholate may be isolated in crystalline form,

either as balls or rosettes of fine needles or in the form of prisms having ordinarily four or six sides (Fig. 99). The bile salts are readily soluble in water. The free acids are slightly soluble in water but readily soluble in alcohol.

The bilo acids have the property of combining with fatty acids to form compounds which have been called choleic acids These are coordination compounds. molecules being joined through secondary valences. Fatty acids containing 16 or more carbon atoms combine with 8 molecules of bile acid; the lower fatty acids, with



FIG. 99 BILE SALTS

from 6 to 1 molecules. These compounds are soluble and diffusible in alkaline or slightly acid solution. For this reason, and because they markedly lower the surface tension in solution and thus promote emulsification, the bile acids greatly assist in the digestion and absorption of fat in the intestines. Through similar combinations they likewise assist in the absorption of ebolesterol, fat-soluble vitamins, carotene, and other substances. They are also responsible for holding the cholesterol of the hile in solution.

There is considerable evidence that the bile acids are used over and over again by the hody. After secretion into the intestine, that fraction of the bile salts which is reabsorbed as choleic acid complexes is liherated in the intestinal mucosa. The bile salts set free, as well as any which havo been absorbed as such, are carried to the liver in the portal circulation. where they become available for resecretion in the bile. This has been called the enterohepatic circulation of the bile salts. It is presumably responsible in part for the marked stimulatory (choleretic) power of the bile salts themselves over bile flow from the liver, since when the bilo salts reach the liver they provide that organ with a readily available supply of a major constituent of the bile itself.

Dila Diamenta The nigments of permel hile are hilimbin and Lat

as ox bile is due to biliverdin. Other types of hile like human hile commonly show the yellowish-brown color of hibrubin (Fig. 100) Modifications of these pigments may be found in gallstones or in altered hile

The bile pigments result mainly from the breakdown of the hemoglobin of the red cells, a consideration of the structures of the hile pigments (p 413) indicates that they are derived specifically from the heme portion of hemoglohin (see Chapter 22) Other heme derivatives in the body-such as catalase, the eytochromes, ete-may contribute to a certain extent to bile pigment formation. The formation of hile pigment has been shown to he a property of the cells of the reticuloendothelial system Since these include the Kupffer cells lining the intralobular

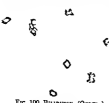


FIG 100 BILIRUBIN (OCDEN)

capillaries of the liver, a part of the bile pigments is produced in the liver Other important sources are the spleen, lymph nodes, and hone mar ron Wherever extravasation of blood occurs, e g , following hrunes, a conversion of blood hemoglobin to bile pigment slowly takes place This con version is apparently going on even in normal red cells, since small amounts of hile pigment have been isolated from this source The first step in the formation of

hile pigments appears to involve an oxidative sersion of the porphyrin ring, to produce carbon dioxide and an open ring compound. In this process the iron is not necessarily lost from the molecule, nor does the open ring compound lose its affinity for globin One such compound of globin and an open ring iron porphyrin derivative is called choleglobin, by analogy to bemoglobin, and other

compounds analogous to those found in the field of heme chemistry are known In general, these pigments are green in color (hence the names verdohemin, verdohemoglobin, verdohemochromogen, etc.) and the open ing of the porphyrin ring has apparently rendered the iron labile, so that it is easily split off by such means as treatment with dilute acid Further steps in bile-pigment formation are obscure, apparently the iron and the globin become detached in some way to produce bilis erdin, which may then be reduced to form bilirubin

Further stages in the transformation of bile pigments involve the action of intestinal bacteria In the intestinal lumen, bilirubin undergoes reduction by bacterial action to form the substance mesobilirubinogen This compound ordinarily is further reduced to form stercobilinogen, which on oxidation becomes converted into stereobilin, the principal pigment of normal feces Stereobilin has been produced in the laborators from mesobilirubinogen by incubation of the latter with normal feces, or with bile-free feces plus added bile. A portion of the stercobilinogen and stereobilin is apparently absorbed from the intestinal tract and reexcreted by the liver thus some gets into the blood stream and ulti matchy is excreted by way of the kidneys into the urine In the urine these two pigments are known respectively as urobilingen and urobilin Considerable confusion has resulted from the nomenclature of these compounds, at the present time it appears to be largely a question of different names for the same substances. Normally the amounts of urobilingen and urobilin in urine are rather small and obviously depend upon such factors as the rate of pigment production and reabsorption, and upon the efficiency of the liver in excreting the reabsorbed material. Thus the urobilingen content of the urine will be low in simple anemias, but high in diseases associated with extensive blood destruction and where liver function is impaired.

The ehemical relationship between certain of the various bile pigments and their derivatives is illustrated by the following structures (Me = CH_1 , Et = CH_2 , X = CH_3 , CH_3).

Bilirubin is insoluble in water but its alkali salts are soluble. It is soluble in chloroform and somewhat in alcohol, but very slightly in ether or benzene. Biliverdin is insoluble in water, ether, or chloroform, but soluble in alcohol. The calcium salts of the pigments are insoluble in water. Solutions of bilirubin exhibit specific light absorption only at the extreme

blue end of the spectrum where the bind merges into the end of the visible spectrum and is not ardinarily detectable as a district bind in the spectroscope. If an iomnomical solution of bilirubin alkali in water is treated with a solution of race chlorid, however, it shows an absorption spectrum similar to that of bilicuanin. When bilirubin is treated with diazotized sulfamilic acid, this regent reacts with bilirubin to form a deeply colored azo dye, known as arobilirubin or arouthin. This reaction is the basis for the Van den Bergh test for line pigments and is allowed in the quantitative determination of bilirubin in biological fluids (see Chapter 23).

Billary Calcult (Billary Concretions, Gallstones), Biliary calcult otherwise designated as bilines concretions or gillstones are frequently formed in the guilbladder. These deposits may be divided into five cla. (1) cholesterol calcula (2) cholesterol calcunor calcula (3) cholesterol edenim pigment calculi (1) edenim pigment edenli and (5) edenh made up almost entirely of morganic material. This list class of calculus is formed principally of the carbonate and phosplints of calcium and is rarely found in man although quite comoron in cittle. The calcium pigment calculus is also found in cattle, but is almost as rare in man as the in organic calculus. This calcium pigment calculus ordinarily consists prin cipally of bilirubin in combination with calcium biliverdin is sometimes present in small amount. The cholesterol calculus is the one found most frequently in min Such calculi may be formed almost entirely of choleterol in which event their color is very light or they may contain more or less pigment and inorganic matter mixed with the cholesterol which tend to give calcult of various colors

Our knowledge of the origin of gallstones is imperfect. Among the forests concerned may be stagnation of bile disturbances of metabolism and infection Chloesterol is held in solution in bile by means of bile neid if cholesterol excretion is increased or bile salt exerction diminished cholesterol stones may form Prolonged stagnation of bile may work in the same direction if the bile salt concentration is reduced by their resorption from the bladder. The resorption of nikaline bile salts touds also to decrease the pill of the bile and thus to reduce the solvent action of the bile salts on cholesterol which is greater to alkaline than in acid solutions. Infection with or without stinguation provides abundant nuclei for stone formation and chemical alterations in the bile far or the formation of cholesterol calcium pigment stones. The impairment in the concentrating power of the mucosa due to noflammation through failure to keep the bile salt concentration sufficiently high may also be a factor.

Okey¹ reports the finding of gallstones in guinea pigs being used for the study of effects of high cholesterol diets. The stones occurred only when cholesterol and riboflavin were added to diets containing 20 per eent protein and were not found when the riboflavin concentration was

For a discussion of cholesterol see Chapter 11, Nervous Tissue

¹ Okey I oc Sor Exptl Biol Med 51 349 (1912)

EXPERIMENTS ON BILE2

- 1. Reaction. Test the reaction of fresh ox bile, using sultable indicator paper. What is the approximate pH of bile?
- 2. Nucleoprotein and Bile Acids. Acidlfy 5 ml. of blle with acetic acid, drop by drop. Note the formation of a precipitate of nucleoprotein and bile acids.
- 3. Inorganic Constituents. Evaporate 10 ml. of bile to dryness in an evaporating dish. Fuse the residue with an excess of sodium carbonate-potassium nitrate "fusion mixture." Cool, extract with 10 ml. of water, and add sufficient concentrated nitric acid to make the extract slightly acid. Filter, and test the filtrate for chloride, sulfate, and phosphate (see Chapter 28).
 - 4. Preparation of Bilirubin from Bile, 500 ml, of blie from surgical drainage patients or 50 ml, of post-mortem galibladder bile (diluted two to three times) is allowed to stand for a few hours in the refrigerator, and the supernatant fluid decanted. This is diluted several times with water and 5 per cent barlum chloride solution is added with stirring. If the precipitate of barlum bilirubinate does not flocculate immediately, add a few drops of 10 per cent NaOH. When the precipitate settles, the supernatant fluid is siphoned off. The precipitate is poured on a filter, washed with water on the paper, dried, and pulverized in a mortar. The powder is extracted with warm alcohol, followed by ether and chloroform, and again air-dried. The precipitate is transferred to a 50-ml. centrifuge tube, moistened with 10 per cent sulfuric acid, and washed three times in a little absolute alcohol, centrifuging and pouring off the alcohol each time. The residue is twice treated in a flask with boiling chloroform and filtered, and the chloroform evaporated, avoiding overheating toward the end. The bilirubin so obtained is rubbed into glacial acetic acid and centrifuged, the acid drained off, and the procedure repeated; the residue is air-dried. It is redissolved in boiling chloroform, filtered, and evaporated to dryness. The final product (40 to 100 mg.) Is brick red in color, free from ash, and quite stable.
 - 5. Tests for Bile Pigments. Practically all of these tests for bile pigments are based on the oxidation of the pigment, by a variety of reagents, with the formation of colored derivatives, eg, mesobilirubia (sellow), mesobiliverdin (green to blue), and mesobilicyanin (blue to violet)
 - a GMELIN'S TEST. To about 5 ml. of concentrated nitric acid in a test tube, carefully add 2 to 3 ml. of diluted blie so that the two fluids do not mix. At the point of contact note the various colored rings: green, blue, violet, red, and reddish-yellow. Repeat this test with different dilutions of bile and observe its delicacy.
 - b Rosenbach's Modification of Gmelin's Test. Filter 5 ml. of diluted bile through a small filter paper. Introduce a drop of concentrated nitric acid into the cone of the paper and note the succession of colors as given in Gmelin's test.
 - c VANDEN BERGH TEST To 5 ml. of diluted bile, add 2 ml. of freshly prepared Ehrlich's diazo reagent.4 Gompare with a control on water alone. In this re-

¹ For experiments on the function of bile saits which may be carried out in this connection see Chapter 19, Intestmal Absorption

See Appendix.

See Appendix.

action an azo dye is formed by a coupling reaction of hilirubin with diazotized sulfanilic acid of the Ehrlich reagent. This reaction is the basis for qualitative and quantitative procedures for estimation of bile pigments in blood serum. A very sensitive test for bile pigment in urine is based on its coupling with a litrobrage diazonium. It discuss sulfanase.

6 Tests for Bile Acids

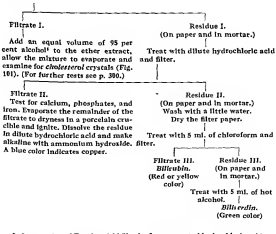
a Furtural-II,SO, Test (Milliss Modification of Pritt Koffes Test) To approximately 5 ml of diluted bile in a test tube add 3 drops of a very dilute (1 1000) aqueous solution of furfural

Now run about 2 to 3 ml of concentrated sulfuric acid carefully down the side of the tube and note the red ring at the point of contact. Upon shaking the



FIG 101 CHOLESTEROL.

- d. Subsoc Tessor Test (Hay). This test is based upon the property that blie acids have of lowering the surface tension of fluids in which they are contained. The test is performed as follows: Cool about 10 ml. of diluted blie in a test tube to 17° C. or lower and sprinkle a little finely pulverized sulfur upon the surface of the fluid. The presence of bile acids is indicated if the sulfur sinks to the bottom of the liquid. Prove this point by repeating the test with water instead of bile. Depending on the grade of sulfur used, this test indicates the lowering of the surface tension from 72 dynes/cm. for distilled water to values below 52-54 dynes/cm.
- 7. Crystallization of Bile Salts. To 25 ml. of undiluted bile in an evaporating dish add enough animal charcoal to form a paste, and evaporate to dryness on a water bath. Remove the residue, grind it in a mortar, and transfer it to a small flask. Add about 50 ml. of absolute alcohol and boll on a water bath for 20 minutes. Fliter, and add ether to the filtrate until there is a slight permanent cloudiness. Cover the vessel and set it aside until crystallization is complete. Examine the crystals under the microscope and compare them with those shown in Fig. 99, Try one of the tests for bile acids upon some of the crystals.
- 8. Analysis of Biliary Calculi. Grind the calculus in a mortar with 10 ml. of ether. Filter.



 Preparation of Taurine. Add 50 ml. of concentrated hydrochloric acid to 150 ml. of bile in a casserole. Boll the mixture in the hood down to a volume

The alcohol is added because it is often found that ery stallization from pure ether does not yield typical cholesterol crystals.

of about 50 ml., filter into a small evaporating dish to remove insoluble material, and concentrate the filtrate by boiling or on the water bath to a volume of ahout 10 ml. Filter the hot solution through a small filter into a 50-ml. graduated cylinder. If the volume is over 12 ml., return the filtrate 50 ml. graduated cylinder. If the volume is over 12 ml., return the filtrate into evaporating dish, continue the evaporation, and filter a second time through the same filter. To the filtrate add 3 volumes of 95 per cent alcoho, mix, and cool in ice water for about 30 minutes. Filter off the crystals on a small funnel, allow to drain thoroughly, then transfer the crystals to a test tube and dissolve by warming in ahout 3 ml. of water. Add 5 volumes of alcohol and allow to stand until maximal crystallization has occurred. Filter



FIG. 102. TAURINE.

Fig. 103. GLYCINE. (SEE ALSO Fig. 38)

off the pure crystals, wash with 5 ml. of alcohol, and allow to dry. Make the following tests upon the taurine just prepared:

- a. Examine under the microscope. Compare with Fig. 102.
- b. Soluhllity in water and alcohoi.
- c. Dissolve a little taurine in water and test this solution for sulfate by adding dilute hydrochloric acid and a few drops of barlum chloride solution is any sulfate present? Boil the solution for a few minutes. Does taurine contain any ethereal sulfate (sulfate hydrolyzable by acid)? To another portion of taurine solution add sodium hydroxide, a few drops of lead acetate, and boil. Is any "lead blackening" sulfur present?
- d. Fuse a little taurine in a porcelain crucible with sodium carbonate-potassium nitrate "fusion mixture." When the crucible contents are color-less, cool, dissolve carefully by the addition of dilute hydrochloric acid, filter into a test tube, and add barium chloride solution. What happens? Explainwhat do these experiments indicate concerning the nature of the sulfur in taurine?
 - 10. Preparation of Glycine. Concentrate the first alcoholic filtrate from Every a until no more alcohol remains. The glycine is present here in the form of a hydrochloride and may be liberated from this combination by the addition of freshly precipitated lead hydroxide or by lead hydroxide solution. Remove the lead from the filtrate by treatment with hydrogen sulfide. Filter

See Appendix.

and decolorize the filtrate by animal charcoal Filter again, concentrate the filtrate, and set aside for crystallization. Glycine separates as coloriess crystals (Fig. 103). See also pp. 131 and 132.

CHEMICAL EVALUATION OF THE FUNCTIONS OF THE LIVER

Many of the metabolic activities of the body are centered in the liver Before discussing certain of these functions and the means for studying them in relation to liver discuse, a resume of the anatomy of the liver seems appropriate. It is necessary also to describe the major diseases of the liver in order to define more clearly the excumstances in which various chemical studies may be made and the response which may be expected

The hver of a healthy adult may weight between 850 and 2600 g, with an average value of about 1600 g Grossly, the liver consists of lobes varying in size and differing to some extent in the sources of afferent blood supplies. Each lobe contains a treelike framework of connective tissue which supports the parenchymal cells blood vessels, and bile ducts. In its finer structure the liver consists of lobules, 1 to 2 mm in diameter, composed of many secretory tubules formed by parenchymal cells surrounding a bile canaliculus. Blood reaches the lobules from the portal vcin and the hepatic artery. The blood enters the sinusoids, spaces between cords or sheets of bepatic cells, and passes through them to drain into the central vein of the lobule.

The pareuchymal cells, the most abundant and characteristic cells in the liver are arranged so that each cell faces both a duct and a smusoid containing blood, and have secretory and metabolic functions. Furthermore the secretory functions may be both execrine and endocrine. In addition to the parenchymal cells, the liver contains large numbers of reticuloendotibelial (Kupffer) cells which form a lining for the sinusoids. The Kinpffer cells have phagoes the properties and are important in connection with hemoglobin breakdown and perhaps in immunity reactions.

DISEASES OF THE LIVER AND BILIARY TRACE

Disturbances of metabolism occurring in liver disease are largely the result of failure of the parenchymal cells to carry out vital functions, because of (1) infections or novious agents, (2) decreased mass of functioning cells, (3) decreased blood supply, (4) impaired nutrition, and (5) reaction of other organs to liver damage, e.g., brain, kidney, pancreas, adrenal, gonads, and splien

Infectious disease of the liver, such as viral hepatitis, is characterized by degeneration and necrosis of parenely mat cells, and may be followed by complete disappearance of cells and destruction of the normal architecture of the lobule. However, regeneration of the parenchymal cells can occur rapidly, and may produce an astonishingly large mass of cells within as little as 24 to 48 hours. Formerly, viral hepatitis was often mistaken for a disease of the bile duets and the term "catarrh il jaundice" in the older literature is a result of this confusion.

Deposition of fat in the liver may occur from overnutrition, from dictary deficiencies, or from the action of toxic substances. Though fatty

liver resulting from overnutration has little importance, deposition of fat caused by toxic substances is evidence of a serious disturbance of hepatocellular function Among the nutritional deficiencies associated with fatty liver are those related to lack of betaine, choline, methionine, or other extrinsic sources of methyl groups Low protein intake may also contribute to the production of fatty livers Liver disease of purely nutritional origin is uncommon in the United States

Liver damage may occur as a result of severe strains on mctabolism associated with various other diseases, such as infectious mononucleosis malaria, lobar pneumonia, typhoid fever, various anemias, syphilis

cholera diabetes, and thyrotoxicosis

Proliferation of the connective tissue of the liver may be of infectious nutritional toxic, hypoxic, or neoplastic etiology, or it may occur spon taneously because of diminished blood supply resulting from circulatory and other factors The overgrowth of connective tissue in turn leads to disorganization of the liver structure and this again to further interference with the blood supply The end result is a shrunken liver consisting largely of connective tissue and with a markedly decreased mass of parenchymal reticuloendothelial, and vascular tissue. The designations portal cirrhosis atrophie cirrhosis and Laennec's cirrhosis have been used at various times to denote such scarred livers, although not all searred livers con form to the pathologists' definition of Laennec's cirrhosis, which accounts for a large proportion of the liver disease seen in American hospitals. A less common form of cirrhosis, known as biliary cirrhosis, is found after prolonged biliary obstruction

Obstruction of the bile duets often eauses jaundice which may be attributed erroneously to liver disease. It is essential to distinguish jaundice due to biliary obstruction from that caused by liver disease or excessive destruction of blood Gallstones entering the common bile duct are the usual cause of bihary obstruction Other causes include neoplastic disease of the ducts and carcinoma of the head of the pancreas Stricture of the ducts may follow infection surgical exploration, or other trauma Disease of the gall bladder frequently is complicated by liver damage The pancreas often shows evidence of being involved Failure to establish the presence of biliary obstruction and to correct it may lead eventually to

biliary cirrhosis

Damage to the liver may be caused by a large number of chemicals and drugs Carbon tetrachloride bas been extensively used for production of experimental liver damage and probably has been involved more often than realized as an insidious cause of clinical liver disease. Atophan various sulfonamides p-aminobeozoate, testosterone, arsenicals, and other drugs have been implicated The effect on liver function, as measured by laboratory studies resembles that observed in biliary obstruction rather than parenchymal liver disease

I iver disease may affect the metabolism and functions of other organs notably the brain and the kidneys Impairment of kidney function com monly accompanies liver disease and may become a grave problem. The coexistence of licpatic and renal failure is often referred to as the hepato-

renal syndrome

THE CHEMICAL PHYSIOLOGY OF LIVER DISEASE?

Carbohydrate Metabolism and Liver Disease. The vital importance of the liver for maintenance of the blood glucose concentration is well established Although hypoglycemia is not a common complication in patients suffering from acute parenchymal liver disease, it occurs in cirrhosis of the Laennee type with sufficient frequency to require that fasting blood glucose determinations be included in the study of such natients Blood sugar concentrations as low as 25 mg per 100 ml are not uncommon Glucose administered to patients with liver disease often causes a greater and more persistent rise in blood glucose than it does in healthy individuals, however, these findings are not of sufficient consistency to permit application as diagnostic or functional tests

Decreased utilization of galactose in liver disease has provided the hasis for one of the earlier tests of liver function 8 The measurement of galactose excretion in urine originally used has been replaced by meas urement of blood galactose concentrations An intravenous galactosetolerance test has also been described, 10 by which the quantity of galactose removed per minute was found to be markedly decreased in liver disease

Serum Cholesterol and Lipides in Liver Disease. The liver is the principal organ concerned with the metabolism and excretion of cholesterol The scrum lipides often show marked changes in diseases of the liver and biliary tract Although serum cholesterol concentration and the partition between free and esterified cholesterol (see p 580) is most frequently studied, neutral fat and especially phospholipide may also show marked changes Relationships between free and esterified cholesterol and between free cholesterol and phospholipide which in health are maintained within narrow limits are subject to striking disturbances in severe parenchymal liver diseaso where lipide concentrations often fall below the minimal levels observed in normal individuals of the same age In viral hepatitis both the concentration of esterified cholesterol and the percentage of the total cholesterol esternied are lowered, if hiver damage is severe esterified choicsterol may become undetectable Recovery is accompanied by rising concentrations of the esterified cholesterol in scrum

Cirrhosis of the Lacanec type is also characterized by low serum lipide concentrations, especially when strophy of the liver is extensive Scrum cholesterol concentrations of less than 100 mg per 100 ml arc quite common The proportion of estcrified cholesterol nlso is lowered, although exceptions exist Low phospholipide concentrations also are the rule

Biliary obstruction regardless of cause is characterized by clevated concentrations of serum lipides Extremely high concentrations, among the highest known to occur due to any eause, are encountered in nationts with biliary obstruction of long duration or with biliary cirrhosis it The

⁷ For a detailed review see Euisely M. II. in Trans. 10th Conference on Later Injury ed teel by F. M. Hoffbauer. New York Josush Maco. Jr. Foundation. 1951.

*Bauer. Hien. Ved. Word. 56. 2538 (1997).

*Parev. Hill. and Nesbitt. J. Lob. Clin. Wed. 35. 705 (1950).

*Colcher. Latek. and Lendall. J. Clin. Invest. 27. 768 (1946).

*Altrens. et al. Medicin. 25. 299 (1950).

reason is not known, since the bile does not appear to be an important route of lipide excretion

Some use is made of serum cholesterol analyses for differentiation of primarily parenchymal lesions from primarily biliary lesions in jumideed patients. However, such analyses offer little information that cannot be obtained more easily and dependably by other methods. The finding of low concentrations and ratios of esterified to total cholesterol in a jain diced patient is strong but not conclusive evidence for parenchymal liver involvement of severe degree. Esterified cholesterol measurements have their greatest usefulness where the more sensitive tests become maximally positive and there is still a substantial amount of functioning parenchymal tissue remaining. As an indication for determination of esterified cholesterol, the presence of jaundice or of a marked elevation of serum hillrubin serves reasonably well.

Information concerning hile acids in liver disease is scant; and unsatisfactory. Improvements in sensitivity of the methods for hile acid determination in serum and hile should overcome this deficiency. Methods available in the past for estimating bde acid conceotrations in serum indicated that hile acid enters the blood stream to attain coocentrations of 10 to 20 mg per 100 ml in the presence of hiliary obstruction. Elevated values may occur also in liver disease affecting the parenchyma predominantly 12

Nitrogen Metabolism in Liver Disease The metabolic transforma tion of amino acids in the liver by synthesis, transamination, etc may be impaired by disease or iojury to produce an abnormal pattern of amino acid content in the blood and urine 12 Certain testa employed for the study of liver function have as their hasis the impaired metabolism of amino acids, for example the tyrosine-tolerance test of Bernhart and Schneider,14 and in its possible relation to gly cine synthesis, the well known hippuric acid test of Quick (see p 915) may also he included bere Amino acids contribute to the characteristic elevation of blood non protein nitrogen content found in severe liver disease along with ures creatine creatinine urie acid and ammonia although the elevated blood VP\ is mainly the result of impaired kidney function. Blood urea mar not be clevated in proportion to creatinine and total \P\ levels, possibly due to impairment of reactions involved in the synthesis of urea Elevated creatine content is not entirely the result of renal failure Interest in blood ammonia in liver disease has been revived by the finding of signs suggesting hepatic come in patients suffering from eirrhosis who had received ammonia-containing ion-exchange resins 13

Plasma Proteins in Liver Disease The liver has a dominant role in plasma protein synthesis being the source of plasma albumin and fibrinogen (and probably other proteins associated with blood clotting) and contributing important components of the a and \(\theta\) globulin fractions. The liver is also involved in the synthesis of \(\theta\) eloquinins, although

there is much evidence indicating that \(\gamma \)-globulin synthesis is largely

The serum albumin level is lowered in cirrhosis, in viral hepatitis during its clinically active stages, in nutritional liver disease, and in neoplastic disease involving the liver Many consider determination of serum albumin to be among the most dependable measurements available for establishing the presence of liver disease (numerous other causes of low albumin concentration usually can be excluded without difficulty) and for following its clinical course For this purpose it is superior to total serum protein concentration because changes in albumin are commonly masked by an equal and simultaneous rise in globulin, so that total protein may remain unchanged or nearly so The lowered concentration of serum albumin is one of the major factors responsible for the occurrence of flocculation in the cephalin-cholesterol flocculation and related tests

An increase in y-globulin accounts for much of the increase in total globulin of serum Whereas in healthy individuals the y-globulin content rarely exceeds 1.6 g per 100 ml, in liver disease concentrations double this are common, and concentrations five or more times the maximal normal occur in some patients with hepatitis. The marked rise in y-globulin occurring in liver discase is similar to that occurring in many other diseases, and may therefore be somewhat nonspecific In many patients it is accompanied by an elevation in the level of β globulin, associated with the appearance of electrophoretically abnormal components in both B and y fractions

Fibrinogen appears to remain within normal limits or to be decreased in liver disease, the a globulin fraction also shows a tendency toward lower levels. This is particularly evident where some of the specific proteins included in this fraction are examined, for example, serum cholinesterase content is markedly decreased in many patients with liver diseaso, as may also be amylase, lipase, and esterase activities

Of the procedures avadable for the study of liver disease which are based upon abnormality in plasma protein production or composition, the determination of plasma albumin and globulin levels are among the most useful These methods are described in pp 601 to 607 Some patients, especially those with liver damage of moderate degree, do not show significant change in serum albumin or total globulin concentrations The quantitative measurement of \gamma-globulin concentration may offer some advantage in this connection and salting out methods are available for this purpose In addition, widely used methods for detecting changes in scrum proteins of the type occurring in liver disease are the semiempirical flocculation and turbidity tests (see p 595) For information concerning tests of this type and an analysis of their mechanisms. Sufer's review 16 may he consulted

Measurement of Excretory Capacity of the Liver. Many substances, among them a number of dyes, are taken up by the liver and secreted into the bile with great rapidty. It has been found that in liver disease the rate of exerction may be lowered, and because of this the

Sufer Am J Mel 13 730 (1952)

use of certain of these substances has been of great value in the measurement of liver function. Disodium phenoltetrahromophthalein sulfonate (sulfobromophthalem, sodium, brom-ulfnlem), introduced in 1925 by Resential and White, has proved to be superior to numerous other substances tested for this purpose. The procedure for the bromsulfalein test for liver function is given on p 598

Detoxification Reactions of the Liver. Numerous studies have demonstrated that in liver disease unrious reactions associated with detoxification are impaired. Among these are included the synthesis of hippuric acid following administration of benzoate. This is discussed in detail on p 915 An improved test employing p-nminohenzorte and based upon serum analysis rather than on urmary exerction has been described 17 Conjugation of various substances with glucuronic nerd has

also been used as n basis for liver-function tests 15

Bile Pigment Metabolism in Liver Disease. Inundice is such a conspicuous sign of liver damage or bile-duct blockage that it has attracted more than its share of attention, often to the neglect of other and more important aspects of liver or biliary-tract disease. The chemistry and metabolism of the bile pigments is discussed on p 411 Methods for the determination of bile-pigment levels in serim (icterie index, Van den Bergh test, total serum bilirubin) are desembed on pp 590 to 595, along with the interpretation of results in relation to liver disease In general, rising scrum bilirulun levels have unfavorable implications, falling values are characteristic of remission of liver disease or biliary obstruction In the urine, bihrubin is normally present in such low concentrations as to be undetectable by ordinary methods, the presence of detectable amounts of bilirubin in urine is indicative of liver damage or biliary obstruction, particularly in the early stage of disease After the presence of the disease has been established, urine bilirubin tests offer little useful information The detection of bilirubin in urine is described on p 836

Bihrubin excreted into the intestinal canal is reduced in part to form urobilinogen which may then be reahsorbed into the blood and again exercted through the bile and also into the urine Normal urine contains detectable amounts of urobilinogen (see p 813), impaired excretory ability of the liver leads to an increased urinary output unless liver dam age is such that secretion is suppressed, or there is biliary obstruction Increased urobilinogen in the urine is a characteristic finding in parenchymal liver disease In the feces urobilingen (stercobilin) levels are increased in hemolytic ancinia and lowered in biliary obstruction

A summary of the relation between liver disease, bile-pigment metabolism, and other aspects of liver function is found in the accompanying table

Other Chemical Manifestations of Liver Disease Abnormalities in blood clotting in liver disease result from a combination of defects, including diminished prothrombin and accelerator globulin levels and

¹⁷ Deus and Cohen J Chu Inzest 29 1014 (19-50).
¹⁸ Ottenberg Wagrach Bernstein and Ilarr w Arch Biochem 2 63 (1943) Snapper and Saltziana Arch Biochem 21 (1999).

COMPARISON OF LIVER FUNCTION TESTS IN LIVER DISEASE

Bilirubin	Tests for Parenchymal Imolement*	Tests for Biliary Tract Involvement
May be present in urine, and may in crease slightly in serum Urobitnogen may increase in urine	BSF usually abnormal CCF, TT, TF, one or more may be abnormal	Vormal
Increased in serum and urine Urobi- linogen generally increased in urine and feces but may be absent	Abnormal	Mainly normal but may be abnormal
Increased in serum and urine	CCF, 'TT TF may be normal or ab- normal, A/G usu ally normal	Increased serum phos phatase, lipide Sim ilar to extrahepatic obstruction
Vay or may not be abnormal in serum and urine Urobi knogen variable	BSF abnormal CCF TT, TF abnormal in about ¾ of the cases A/G abnor- mal	Variable Phosphatase may be high
Variable Intermittent or continuous elevation Extreme elevation Very lon fecal uro bilinogen	Generally normal, but liver parenchyma may become in jured and tests posi- tive Same as preceding	Variable with elevated serum phosphatase and lipide preva- lent Good response to vitamin K. Elevated serum phos phatase and lipide Good response to vitamin K.
Bilirubin elevated Urobilinogen vari able	BSF increased A/G abnormal TT in creased CCF increased in ½ of the cases.	Elevated phosphatase Marked elevation of serum lipide especi ally phospholipide
moderately ele- vated, direct bili	Seldom abnormal but may become so due to hypoxia or other complications	Seldom abnormal Pig ment stones may cause biliary ob- struction
	Variable Internitient or continuous elevation Extreme elevation Variable Internitient or continuous elevation Extreme elevation Extreme elevation Very lon fecal uro bilinogen Extreme elevation Very lon fecal uro bilinogen Bilitubin elevated Urobilinogen Total serum bilitubin moderately elevated Urobilinogen Total serum bilitubin moderately elevated, direct bili urine urobilinogen increased, urine urobilinogen increased Urine bilitubin negative propositiones elevation received Urobilinogen increased, urine urobilinogen	May be present in turne, and may in crease slightly in serium Urobilinogen may increased in urne and urne Urobilinogen generally increased in urne and frees but may be absent Increased in serum and urne urobilinogen generally increased in urne and frees but may be absent Increased in serum and urne urobilinogen generally increased in urne and frees but may be normal or abnormal A/G usually normal Vary or may not be abnormal or abnormal in serum and urne urobilinogen variable Variable Internit tent or continuous elevation Very lon feeal urobilinogen Very lon feeal urobilinogen Very lon feeal urobilinogen Bilirubin elevated Urobilinogen variable Bilirubin elevated Urobilinogen variable Same as preceding BSF mereased A/G abnormal TT in jured and tests positive Same as preceding BSF mereased A/G abnormal TT in ablot ¼ of the cases. Seldom abnormal but may become so due to hypoxa or other complications

^{*}Abbreviations BSF bromsulfalein A serum albumin G serum globulin CCF cephalin-cholesterol flocculation TF, thymol flocculation TT thymol turbidity

probably other factors. The depletion of prothrombin in biliary obstruction is related to impaired vitamin K administration provided liver damage is not excessive. The evaluation of the elotting function of the blood is discussed in Chapter 22.

Liver disease may be associated with changes in blood and urine levels of such substances as citrate, 20 lactate, pyruvate, succinate, etc., but study of such changes has not yet reached the stage of general clinical application. Hormone metabolism may also be altered in liver disease, for example, both the conjugation of injected testosterone²¹ and its exerction as 17-ketosterond²² are decreased in liver disease.

Disturbances of water and electrolyte balance are among the more significant and troublesome complications of liver disease. These are due in part to impairment of liver function and in part to the effect of liver disease on the kidneys and other organs. Among the blood analyses which are of clinical importance under these conditions are those for plasma proteins (p. 601), scrum sodium (p. 619), and scrum potassim (p. 652). Scrum iron levels may also be altered in liver disease, being clevated in hepatitis²² and in hemochromatosis, in the latter condition, the iron binding capacity of the scrum may be at the saturation level as compared with 14 to 69 per cent of capacity in cirrhosis and 28 to 59 per cent of capacity in normal individuals ²⁴

Measurement of the alkaline phosphatase activity of scrum (see p 635) often assists in the differentiation of parenchymal liver disease from that due to obstruction and other lesions of the bihary tract Bihary obstruction is characterized by a persistent increase in phosphatase activity to two or more times the maximum found in healthy individuals. This rise occurs regardless of the nature of the obstruction, whether due to calculus, stricture, or neoplasm Elevation of scrum alkaline phosphatase may provide one of the few clues to the presence of neoplastic growth in the liver.

Elevated values are also found in toxic hepatitis of chemical origin and in inflammatory disease of the hile ducts (cholangiolitic hepatitis) but are uncommon in cirrhosis of the Lacinice type The factors responsible for variation in scrum phosphatase activity in liver disease are not clearly understood

Serum cholinesterase activity (p. 638) is usually lowered in the presence of atrophy of or damage to the liver parenchyma, although not consistently Serial measurement of cholinesterase activity appears to provide a useful method for following the course of hier disease, particularly that associated with deficient diets

Selection of Procedures The choice of methods for the study of disease of the liver or bihary tract will vary according to the type of information sought. The following outline shows the principal purposes for which chemical methods are applied together with the first the pro-

cedures most likely to prove useful A decision regarding the number of tests to be used requires experience and judgment. Usually those tests stalicized will suffice for an initial study. To apply simultaneously the entire group of tests listed in any of the categories would seldom be nistified

1 Detection of liver damage in absence of jaundice, e.g., early or subclinical hepatitis urine bilirubin, bramsulfalein retention, direct and total serum bilirubin, flocculation tests (eg, cephalin-cholesterol flocculation,

thymol turbidity and flacculation), urine urabilinogen

2 Detection of residual liver damage, "recovery" stages of hepatitis, chronic hepatitis, chronic passive congestion, portal cirrhosis bromsulfalein, direct and total serum bilirubin, flocculation tests, serum albumin and globulin, prothrombin, serum cholinesterase, urine urobilinogen, and coproporphyria

3 Following the course of the jaundiced patient suffering from parenchymatous disease, direct and total serum bilirubin, flocculation tests, serum albumin and globulin, prathrombin In addition, if severe, serum

esterrited cholesterol, cholinesterase

4 Differentiation of mundice due to biliary disease from that due to parenchymatous disease serum alkaline phosphatase, cephalin-cholesterol flocculation and thymol turbidity, galactose tolerance, prothrombin, feces urobilinogen

5 Differentiation of extrahepatic biliary obstruction due to calculus

from that due to neoplasm, stricture, etc feces urobilinogen

6 For following the course of the surgical patient with disease of the biliary tract plasma prothrombin, phosphatase, direct and total serum bilirubin, albumin and globulin, electralytes, blood urea N or NPN, serum lipides

7 Differentiation of hemolytic jaundice direct and total serum bilirubin, feces urobilinogen, erythrocyte fragility, reticulocyte count

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19

Intestinal Absorption

Mechanism of Absorption. Absorption is primarily a function of the small intestine Very little absorption takes place from the stomach, and the process is nearly complete before the colon is reached The small intestine is particularly adapted to carrying on this process by virtue of its great length (about 9 meters in man) and by the structure of its mucous membrane The surface of the latter is greatly increased (to a total area of about 10 square meters) by the presence of folds and of fingerlike processes called will The latter possess a rich blood supply, lymph spaces called lacteals, and muscle fibers whose contractions mechanically assist absorption The mucous membrane through which substances must pass to enter the blood or lymph is extremely thin, but it must not be forgotten that it consists of living cells

Substances in solution tend to distribute themselves uniformly throughout the solvent, and this tendency, which we measure as osmotic pressure, is a powerful force Undouhtedly osmosis may play some role in the absorption of certain substances which may be present in the intestine is far higher concentrations than in the blood. There is much evidence

however, that the process is rarely one of simple diffusion

The facts with regard to the absorption of ionic substances (salfs, amino acids, etc.) have been made much more intelligible by the development of the Donnan theory of membrane equilibria. This explains how it is possible for ionic diffusion to be more or less selective and how certain ions may pass into the blood although present there in higher con-

centration than in the intestinal lumen

Even with this extension, however, it is not yet possible to explain many phenomena of absorption on a purely physical basis. Not only do the regulatory powers of the cell and its changing structure influence the course of diffusion processes but it seems probable that certain substances in their passage enter into chemical combination with constituents of the protoplasm thus making possible different types of selective transportation. Furthermore, white blood cells actually migrate back and forth between the lymph and the lumen of the intestine. There is evidence that certain of these may actually engulf particles of iron compounds and fat droplets One of their functions may thus be to assist by phagocytoss in the absorption of difficultly soluble substances. Their number is bardly adequate to permit their playing a large role in the absorption of food stuffs, however, their accumulation in the intestinal microsal during digestion would indicate that their function is an important one

The intestinal mucous membrane further possesses the power of excreting certain substances into the gut. Sucrose injected into the blood stream appears in part in the duodenum and calcium salts are eliminated by the mucosa of the large intestine. The glands of the mucosa secrete a digostive fluid the intestinal juice and the cells themselves contain enzymes such as proteinases and peptidases, carbobydrases and phosphatases which help to prevent the passage of incompletely decomposed proteins, carbobydrates and organic phosphates into the blood.

Because of the many factors involved, the study of intestinal absorption offers certain difficulties. Investigation of the processes concerned helps however, to throw light on permeability and other fundamental properties of protoplasm. It is further of the greatest practical importance to under stand the factors governing the absorption of the many essential food stuffs and to determine the conditions under which there is a loss of the great protective power of the intestinal mucosa to detoxicate or prevent the absorption of toxic substances which may be ingested or produced during digestion or putrefaction.

Mineral oil has been shown to interfere with the absorption of fat soluble vitamins and various digestion products Certain adsorbing agents used therapeutically also interfere and all catharties interfere because of the reduced time the products of digestion remain in the

intestine

Absorption of Carbohydrates Carhohydrates are absorbed almost entirely as the simple monosaccharides glucose fructose galactose mannose and the pentoses each having a characteristic absorption rate Corn showed that galactose was absorbed most rapidly Next came glucose and fructose L-Arabinose bad the slowest absorption rate less than onetenth that of galactose and glucose There is considerable evidence favor ing the view that a hevolinase-catalyzed phosphorylation takes place in the intestinal absorption of glucose followed by a dephosphorylation of the sugar by phosphatases in the mtestine before it enters the circulation Of the disaccharides lactose is least readily by drolyzed in the intestine and some of it occasionally reaches the blood as such. In these cases it is eliminated unchanged in the urine Sucrose entering the blood is also excreted as such When strong sugar solutions enter the bowel they are diluted by the intestinal secretion until the concentration is reduced to a point favorable for absorption 1 e when the osmotic concentration equals that of the blood plasma Lactose, because of its slow digestion and absorption reaches a much lower portion of the bowel than other sugars and is thus more effective in promoting the growth of neidophilic rather than putrefactive bacteria

The simple sugars entering the blood are quite rapidly removed therefrom by the liver and other tissues for oxidative and storage purposes. After the ingestion of 100 g of glucose a maximum blood-sugar value of about 0 15 per cent is reached in less than an hour the normal figure of about 0 1 per cent being restored by the end of two hours. This rapid absorption of glucose and its subsequent oxidation in the tissues to yield energy are of importance to persons engaged in prolonged athletic competition or in other fatiguing activities.

If the intestinal inucous membrane is destroyed or poisoned by such i substance as sodium redorectate, it nits like nu ordinary permeal!e membrane and sugars pass through it according to ordinary laws of diffusion. Thus the pentos is pass through more rapidly than the hexages With the normal living membrane however, the hexosesshow more rap d ab orption. Apparently the hexo es are combined with phosphate is pasing through the cells of the mucous membrane. The formation of the compound greath speeds up the absorption process and makes possible absorption even from a sugar solution of lower concentration than the blood. The intervention of chemical mechanisms of the hyung cell also makes possible a certain selectivity in absorption, and an adaptation of the mucous membrine in the direction of self-protection of the body against rapid absorption of toxic substances

Absorption of Protein Protein is absorbed for the most part in the form of individual amino acids. The mains need content of both portal and systemic blood rises after a meal. The amino acids of the di t enter ing the body are carried by the blood to all the tissues of the body, where they rapidly become incorporated into the metabolic processes of the cells and become indistinguishable from the namio acids already present unless they have been labeled in some way, as with isotopic introgen Thus the dutary ammo acids are not to be regarded as somewhat of a surplus as they were at one time, to be drawn upo i or oxidized as the need arose but rather as a daily contribution to the general processes of

nitrogen metabolism in the body (see Chapter 33)

Whereas proteins are absorbed mainly as minio acids it is nevertheless true that certain molecules larger than the amun acid molecules eg peptides or polypeptides may at times be abserbed Strepogenin peptidelike substance is definitely absorbed With the aid of immunologi cal methods it has also been established that certain unsplit proteins may be absorbed unchanged

A variable and sometimes significant proportion of the dietary amino acids escape absorption and are metabolized by the bacteria of the intetinal tract. The products of this action may be absorbed and appear in the blood or urine The relatively high ammonia content of portal blood is attributed to these processes as is the indican content of the urine The significance of such intestinal putrefaction is discussed in Chapter 20

Incompletely decomposed protein may sometimes be absorbed, as 15 indicated by the production in certain individuals of anaphylactic reactions (cutancous eruptions asthma etc.) following the ingestion of par ticular types of protein Such reactions are not produced by completely hydrolyzed proteins

Absorption of Fat The exact mechanism by which fat is absorbed is the subject of considerable controversy. Two theories have been proposed namely the lipolytic and the partition theory and each of thee possesses certain ments According to the hipolytic theory as defined by Bloor as well as by Verzar a complete hydrolysis of the ingested fat occurs in the gastrointestinal tract as a preliminary to absorption Glycerol formed as a result of such hydrolysis is readily soluble in water,

Woolley Pederation Proc 6 424 (1947)

there is considerable experimental evidence that this compound can be readily and completely absorbed. On the other hand, the fatty acids resulting from hydrolysis of the neutral fats are insoluble in water, although their somes dissolve quite readily. However, the pll of the small intestine is such as to preclude the presence of appreciable amounts of soap. Since it is known that most fats are almost completely digested, some mechanism is required to transport the water-insoluble fatty acids through the gut wall if soaps are excluded. The mechanism seems to involve the presence of bile concomitant with the fat. Under such conditions an emulsion is formed which is reflected in the milky appearance (chyle) of the lymph in the mesenteric vessels during the absorption of fat. This condition does not obtain in the phsence of bile. Bile is believed to facilitate digestion both by activating panerentic lipase and also by rendering the fatty acids soluble in the aqueous medium. The latter change is mediated by the bile salts, which lower the surface tension, It has been postulated that the bile salts combine with the fatty avids to form compounds known as cholcic acids. Wieland and others have demonstrated that such compounds, produced in the test tube, have definite melting points, and that there are fixed ratios of fatty acids to bile acids (usually 1:8 in the case of Cie-Cia acids). The importance of this mechanism remains questionable, however, since the only unturally occurring bile acid which is active is deexycholic acid, and this is a minor component in most biles. Moreover, only the unconjugated form is active; but the conjugated acids (glycodeoxycholic and tanrodeoxycholic acids) represent the forms occurring in the bile. The fact that such coordination compounds as cholcie acids exist is however an intriguing one, and theso compounds may be related to fat absorption.

According to the partition theory, as proposed by Frazer, fats are only partially hydrolyzed in the gut, with the resultant formation of some fatty acids, and also mono- and digtycerides. The bulk of the fat is in the form of triglycerides. These lower glycerides and fatty neids, togethor with bile salts, form a stable emulsion which consists largely of triglyceride fats. This particulate matter, when the droplets are of a size of 0.5 ar less, is directly absorbable through the fine canals in the outer border of the intestinal cells. The results of Mattson et al. (see Chapter 33) tend to support this hypothesis. They showed that the intestinal contents of rats, removed three hours after the animals had been fed a mixture of partially hydrogenated cottonseed and soybean oils, contained principally unjudyced triglycerides, diglycerides, and monoglyrerides, with only 15 per cent of free fatty acids. Frazer is of the opinion that triglycerides of short-chain fatty acids, whose acids are water-soluble, may be completely hydrolyzed in the intestine.

There are ample experimental data which demonstrate that, irrespective of the fatty material fed, neutral fat appears in the intestinal lymph. According to the Verzár school, the products of fat digestion are recombined in the intestinal mucosa, possibly through the intermediation of plospholipide. Frazer also suggests that considerable medification and resynthesis of triglycerides may occur in the intestinal rells. The bile acids set free are returned to the lumen of the gut to assist in the absorption of additional fat. Atthough new phospholipides may be formed in the intestinal cells, as demonstrated by the incorporation of the un natural fatty acid elaidie acid in the intestinal phospholipides after the feeding of trielaidin (Sinclair), considerable quantitative data are non available which indicate that the phospholipides are not obligator, intermediates in fat synthesis

By means of tests (rats) in which decanoic and palmitic acids labeled with C14 in the carboxyl position were fed, it was learned that shortchain fatty acids are transported mainly by the portal pathway, and

long-chain saturated fatty acids via lymph 2

After resynthesis, the greater part of the fat first enters the lacteals of the intestinal villi and then the lymphaties, forming an emulsion (the chyle) which is carried via the thoracie duct to the jugular vein An in crease in the fat content of the blood and lymph following a meal ("fat tide") is readily demonstrated, the fat appearing in the form of minute globules called chylomicrons The fat of the diet is therefore unique among the other components of the diet in that in large part it bypasses the liver The significance of this anatomical arrangement is not clear, but it may possibly be related to the recognized ability of liver tissue to oxidize fatty acids rapidly to the stage of acetoacetic acid and β-hydroxybutyne acıd

Absorption of Sterols. Cholesterol, the most important of the sterols, like the fatty acids forms compounds with the bile acids which facilitate its absorption (see Chapter 18) Unabsorbed cholesterol 18 reduced to coprosterol It is uncertain whether this change to coprosterol is brought about solely through bacterial action. The plant sterol pbytos terol is not absorbed and hence cannot become a source of cholesterol in the animal body Ergosterol is said to be absorbed but slightly if at all, but the irradiated form (ealciferol) is more readily absorbed Apparently the absorption of sterols is very specific, so that even isomerism may alter absorbahility, and saturation of unsaturated bonds may change a stero from a readily absorbable substance to one completely unabsorbable Of the many sterols found in plant or animal foods, the only one which is absorbed in the human intestine, aside from the D vitamins, is cholesterol Certain forms of vitamin D are absorbed to different degrees in different species (see Chapter 35)

Absorption of Inorganic Salts. The selective nature of absorption applies even to inorganie salts. Thus sulfates are much less readily absorbed than chlorides and tend to withdraw water from the blood For this and other reasons the sulfates have a cathartic effect. The absorption of calcium and phosphorus is of especial interest because of its relation to the development of rickets and because this absorption can be so profoundly affected by minute amounts of antirachitic vitamin and hence by ultraviolet radiation The acidity of the intestinal contents is also of importance for the absorption of the relatively insoluble salts of calcium The most rapid rate of absorption of calcium after the administration of CaCl, solution containing Cass has been found to occur within the first 2 to 4 hours This absorption was principally from the proximal part of the small intestine Dietary iron appears to be absorbed in nutritionally

Kiyasu Bloom and Charkoff J Biol Chem 199 415 (1972) Harrison and Harrison J Biol Chem 188 83 (1951)

significant amounts only when it is in the ionized inorganic form, and clinical and experimental evidence indicates that ferrous iron is much more available for absorption than ferric iron. Calcium, magnesium, phosphorus, and iron are excreted to a considerable extent by the intestinal mucosa. Hence a study of their absorption at different levels of the intestinal tract is required for an understanding of the factors involved. The absorptive power of the colon for food substances is relatively low.

Methods of Studying Absorption. Mucb of our information on absorption has been obtained from the study of isolated intestinal loops which retain their nerve and blood supply. Solutions can be injected into such loops and the contents removed at any time for analysis. The relative rates of absorption of various substances can thus be determined under controlled conditions Small animals such as the rat may be sacrificed after the ingestion of test substances, the intestinal tract removed, washed out, and the washings then analyzed to establish the extent of absorption.

Histological examination of the mucosa of animals killed after a meal has shown the presence of fat globules in the cells.

By establishing fistulas of the intestine, the course of absorption in different parts of the tract has been studied.

Analysis of the blood and lympb gives information of great importance relative to the nature of the products entering the blood stream Urine analyses show the rapidity with which soluble inorganic salts are absorbed. Perfusion experiments on the intestine are little used because a normal mucous membrane is difficult to maintain. Studies on absorption from the stomach may be made with a stomach tube, and a similar procedure may give some information as to absorption from the colon. Fecal analyses show the completeness of digestion and absorption of various substances present in the food.

Animals may also be fed a diet containing a definite amount of a nonabsorbable substance such as iron oxide, or a labeled component such as claidic acid or compounds containing the isotopes of hydrogen, phosphorus, nitrogen, iron, or carbon By suitable analyses of the intestinal contents and the other parts of the animal body the rate and extent of absorption may be evaluated. The use of isotopes in particular appears to offer great promise in clucidating the mechanism of absorption. For a discussion of isotopes, see Chapter 32, p. 970.

EXPERIMENTS ON ABSORPTION

- 1. Experiment to Show the Action of Bile Solt Solutions on Fatty Acids and Cholesterol. Prepare five test tubes as follows:
- a. Five ml. of buffer solution pH 7 + 2 ml. of a 10 per cent solution of bile salts + 1 ml. of a 1 per cent solution of oleic acid in alcohol.
- b. Five ml. of buffer solution pH 7+2 ml. of water +1 ml. of 1 per cent oleic acid solution.
- c. Five mi. of buffer solution pH 9 + 2 ml. of water + 1 ml. of 1 per cent oleic acid solution.
- d. Five ml. of water + 2 ml. of 10 per cent blie salt solution + 1 ml. of a 0.05 per cent solution of cholesterol.
- e. Five mi. of water + 2 ml. of water + 1 ml. of cholesterol solution.

 Place all tubes in a water bath at 40° C. for a few minutes and observe the

tubes for turbidity.

Fatty acids form a clear soap solution only at pil 9 or higher. With bile salts a clear and diffusible solution is formed even below pil 7. Bile salts also have a similar action on choiesterol. This is important also in connection with the excretion of cholesterol in the bile.

 Determination of Rapidity of Absorption of Sugars from the Entire Gastrointestinal Tract (Method of Cori): Principle. Annuals are given sugar solutions by a stomach tube. After a suitable time has dipod the annuals are hilled, the entire gastrointestinal tract removed, and the total sugar remaining unabsorbed determined.

Procedure. Rats two to three months old and weighing from 120 to 18 g, are weighed. They are then placed in small wire-screen cages with screen bottoms so that there is no access to fecces. For 48 hours they are given water but no food. The rats are again weighed and are then fed the solutions to be tested, usually 1,25 to 2.5 ml, of 25 to 80 per cent sugar solutions. These are introduced by means of stomach tunes consisting of Nos. 4 to 5 urethral catheters softened by plunging for a moment into holling water. A small mouthpiece is used and the catheter marked to indicate the depth to whichlit should be introduced. A syringe of the Record type with a needle to connect with the catheter is used to inject the fluid. If diarrhea is caused the experiment is discarded.

An animal is killed at each hourly interval. The entire gastrointestinal tract is removed, slit open, and washed thoroughly with water making up to a volume of nearly 500 mi. A small amount of dialyzed iron is added and then a little sodium sulfate to precipitate it, alone with interfering substances. The sugar is determined by the Benedict method or some other method. Blood sugar may also he determined by the liagedorn-Jensen method or other micro method. Sugar may also be determined in the urine (in another experiment it may be shown by analysis of the gastric contents that a dilution of the sugar solution takes place in the stomach.) The amount of sugar absorbed per 100 g. of body weight of the animals is calculated. An absorption curve may also be piotted from the results obtained with similar animals at t, 2, 3, 4, etc., hours. Cori found that the rate of absorption of hexoses is independent of the concentration, and the rates of absorption of some monosaccharides are in the following order, galactose > glucose > fructose > mannose > xylose > arabinose. No glucose appeared in the urine but about 50 per cent of the galactose was excreted by this channel.

If a soluble ferric salt such as ferric ammonium citrate is added to the sugar solutions it is possible to determine in what parts of the tract digestion and absorption of carbohydrates, etc., more particularly occur (see the experiment below).

The tolerance of animals for sugars injected intravenously may also be determined and an idea obtained as to the rapidity with which sugars a absorbed when injected intravenously as compared with absorption from the intestine. Corl found the tolerance for glucose given intravenously to be from 2.2 to 2.5 g. per kg of body weight per hour.

3. Influence of Carbohydrates on the Utilization of Calcium and Phosphorus (Method of Bergeim*): Principle To a standard diet is added a definite

Bergeim J Biol Chem. 79, 29 (1926) Gallup J Biol Chem. 76, 43 (1928). Heller Breedlove and Likely J Biol Chem. 79, 275 (1928) For use of alice ace Greenwald and Gross J Biol Chem. 64, 185 (1925). 87 505 (1929) Gallup J Biol Chem. 81, 23 (1929) For use of the iron method in the study of dingestion and absorption in different parts of

proportion of iron oxide and of the carbohydrate whose effect it is desired to study The ratios of calcium and phosphorus to iron are determined for foods and feces and the percentage absorption calculated The accurate separation of feces of the evperi mental period is not necessary

Procedure Feed two or more albino rats (about 60 g in weight) on a phos phorus low and calcium high dlet (whole yellow corn 76, wheat gluten 20 calcium carbonate 3 sodium chloride 1, and c p ferric oxide 0 2) for about three weeks. Put in separate cages with screen bottoms. Collect feces for a five day period. Then modify the diet of the animals by substituting chicase for 30 parts of corn in one case and an equal amount of factose in the other After a two day interval collect the feces for a period of four days. Then ex change diets so that the first animal dets lactore and the second clucose After a two day interval collect feces ngaln over a four day period. All feces need not be collected. Those contaminated with urine are discarded

About 0.5 g (not weighed) of feces from each collection is ashed, preferably in a 35 ml silica crucible at a moderate temperature. Add a few drops of nitric acid to the residue and heat again to destroy the last carbon Add 10 ml of 15 per cent hydrochloric acid and heat until the ash is dissolved Wash into a flask with water to make about 35 ml Ash 2 g of food and dissolve the ash in the same way Determine iron, calcium, and phosphorus in the ash solutions by standard methods (see Chapters 23 and 31) and record the num ber of me of each in 1 ml of ash solution

CALCULATION Calculate the ratios Ca/Fe and P/Fe for food and feces and calculate percentage absorption or utilization of Ca and P. For example of the ratio Ca/Fe for food is 10 1 and in feces 4 1 unabsorbed Ca is 4/10 or 40 per cent, and utilization is 100 - 40 - 60 per cent

Lactore promotes calcium absorption by creating an acid medium (lactic acid) in the intestines Glucose has little effect Vitamin D markedly improves absorp tion of Ca and P

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^{3 110 (19%)}

Putrefaction, Detoxication, and Conjugation

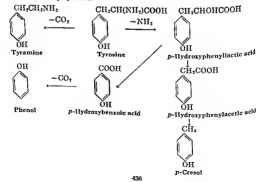
The food residue and the digestive and other secretions as they pass into the lower leum and colon are acted upon by the bacterial flora which become permanently established in man during the first few days of life The extent of bacterial activity is indicated by the fact that nearly one-

third of the solid matter of normal feces is made up of bacteria.

The metaholic products of the intestinal bacteria are for the most part harmless, and some are actually useful; but a few, especially those aresing from proteins and amino acids, are toxic in varying degrees. In the past it was believed that an excess production of these products, such as might occur in constipation, caused a condition of autointoxication characterized by malaise, headache, and irritability. Probably mechanical factors as well as emotional and psychic influences often account for some of these disagreeable effects attributed to constipation.

When amino acids are acted upon by bacterial enzymes, they undergo decarboxylation, dearmination, and other reactions, with the result that abnormal products are formed. Some of these are toxic and are handled in the body by chemical mechanisms that are generally regarded a detoxifying The variety of products that may result from an amino acid

is illustrated by tyrosine.



Action of Products of Putrefaction. Of the various types of compounds formed by bacteria acting on amino acids, the amines are physiologically the most active. Tyramine, which is formed from tyrosine, is a strong vasoconstrictor and like epinephrine, to which it is closely related structurally, it elevates blood pressure. Histamine, on the contrary, causes capillary dilatation. Although this compound may be formed by bacterial action, it is also normally produced by the body to serve as a stimulus for the secretion of hydrochlorie acid in the stomach. In anaphylaetic shock large amounts of histamine are released into the circulation. Since tryptamine is a product of putrefaction, it is interesting to recall that when platelets disintegrate during the clotting reaction, 5-hydroxytryptamine is produced. This compound is a strong vasoconstrictor and probably plays an important role in hemostasis. Of the aliphatic amines, two are well known: pentamethylenediamine (eadaverine) and tetramethylenediamine (putrescine), which are derived from lysine and arginine respectively. Since they were originally isolated from putrefying flesh, they were classed as ptomaines. In spite of their offensive names, they are relatively harmless. The condition known as ptomaine poisoning is due to haeterial toxins and not to these amines

Of the putrefactive products other than amines, indole and skatole, which are derived from tryptophan, have received much attention. Both possess a dasagreeable odor and are mainly responsible for the characteristic odor of feces. Only a small fraction of the indole and skatole formed in the intestines is absorbed, as indicated by the finding that the daily exercition of indican (indoxyl potassium sulfate) rarely exceeds 10 to 20 mg. It is improbable that such small amounts have any deleterious effects Phenol, p-cresol, and allied compounds formed from aromatic amino acids are toxic, but the quantities formed and absorbed are rarely large enough to be harmful provided the liver and kidneys function normally. Total phenol excretion by normal men averages about 0.2 g. ner day.

Some of the bacterial products formed in the intestines are distinctly useful to the body. It is likely that part of the vitamin K required by the organism comes from the action of E. coli and other bacteria. It is known that various other vitamins such as biotin and thiamine are synthesized by bacteria, and that these can be utilized by the body. The bacterial flora may therefore he regarded as an auxiliary factor in nutrition. The benefits thus accruing are likely to outweigh the harm that may come from the toxic putrefactive products.

DETOXICATION

Though it is well known that the body can and does develop means of defense against microorganisms and their toxins, which usually are proteins, there is no convincing proof that it has evolved mechanisms specifically intended to detoxify either the products resulting from putrefaction in the intestines or the myriads of compounds and drugs that are ingested and absorbed.

¹ Rapport, Green, and Page J. Biol. Chem , 180, 961 (1949).

In general all products of exogenous origin that enter the body may be divided into three major classes (1) Substances which are normally utilized for energy structure or function All may be broadly considered as food (2) Substances which in the doses ingested after physiological function If the change in function is beneficial the substance producing it is considered to be a drug whereas if the alteration is harmful the substance is considered to be a poison (3) Substances which pass through the body unaltered and cause little or no change in function Water is the most important example although in a strict sense it may be regarded as a food

In categories as broad as these many instances of overlapping as well as difficulties of interpretation arise Ethyl alcohol may be regarded as a food a drug or a poison Similarly, nicotinic acid being a vitamin 15 a food but it may also act as a drug and in large doses as a poison

Although the protection against noxious substances is probably largely accidental rather than by design detoxication nevertheless plays an important physiological role. It begins in the intestines. Thus, the enzyme histaminase found in the alimentary tract is probably responsible for the destruction of relatively large amounts of histamine. The intestinal wall is coordinated in the line of defense against poisons since it acts as a barrier to various harmful substances Many toxic substances that pass the intestinal walls are removed by the liver. It is not surprising therefore that this organ which has as one of its functions that of a chemical watch dog possesses many of the mechanisms which are usually designated as detoxication reactions The principal reactions namely oxidation reduc tion hydrolysis and conjugation which the body normally employs for the destruction detoxication and elimination of a foreign compound are not essentially different from those employed on normal metabolites Many foreign substances are only partially metabolized thereby leaving a chemical clinker or residuum which often serves as a valuable source of information concerning metabolic pathways

Oxidation, Reduction, and Hydrolysis The first attempt of the body to protect itself against a toxic compound is to destroy it by oxida tion Many substances such as ethyl alcohol are completely burned to earbon dioxide and water Sometimes the intermediary oxidation products are more toxic than the original compound as illustrated by methyl alcohol which yields formaldeby de and formie acid

The oxidation of aromatic compounds is of great physiological impor tance Benzene itself as well as many simple aromatic compounds such as benzoie acid and phenol are fairly resistant to oxidation Certain side chains particularly the groups CH-CH(\H1)COOH which occurs in phenylalannie and CH2CO COOH render the aromatic nucleus com pletely oxidizable This suggests that oxidative deamination is the initial step in the metabolism of aming and

having a side chain with an odd number of earbon atoms yield benzoic acid, and those with an even number yield phenylacetic acid

 $C_6H_5CH_2CH_2CH_4COOH \rightarrow C_6H_6CH_2CH_2COOH \rightarrow C_6H_5COOH$ Phenylvaleric acid Phenylpropionic Benzoic acld acld

 $C_6H_5CH_2CH_4CH_2COOH \rightarrow C_6H_5CH_2COOH + CH_3COOH$ Phenylbutyric Phenylacetle Acetle acid acid acid

From these observations Knoop formulated the hypothesis that normal fatty acids are likewise entabolized by \$\beta\$ oxidation (see Chapter 33)

The finding? that the departreatized dog oxidizes phenylbutyric acid to phenylacetic as readily as a normal dog is interesting because it was the first evidence suggesting that the oudation of butyrie acid was not impaired in diabetes

Reduction and hydrolysis are encountered infrequently in the metabolism of foreign compounds. One example of reduction is the conversion of chloral hydrate to trichloroethyl alcohol, which is then combined with glucuronic acid Glucosides such as the digitalis compounds probably undergo hydrolysis with the liberation of a sugar and an aglycone as a step in their metabolism and elimination from the body

The conversion of cyanide to thiocyanate may be regarded as an important detoxication mechanism. This is indicated by the wide distribution in various tissues of the enzyme rhodanese, which is responsible for changing cyanides to thiocyanate, and also by the relatively high concentration of thiocyanates in the blood (over 1 mg per 100 ml) The reaction may be expressed as

The source of the sulfur is not known. It can be supplied by thiosul fate in titro but it remains uncertain whether this compound is utilized

physiologically

Conjugation. The isolation by Keller in 1842 of hippuric acid after the ingestion of benzoic acid led to the recognition of a new type of biochemical reaction, namely, the conjugation of a foreign organic compound with a normal metabolic product such as glycine Other compounds or radicals employed by the organism for conjugation are glucuronic acid, sulfuric acid, cysteine, glutamine, ormthine acctic acid, and the methyl group

Antimetabolites Though it is generally assumed that conjugated products are the resultants of detoxication at is more likely that they are merely the end products of normal metabolic processes applied to foreign compounds When any ingested compound is excreted in a conjugated form it may be regarded as an antimetabolite (see Chapter 36). One may postulate that such a compound has enough structural similarity to a normal metabolic product to engage the enzyme designed for the latter

² Quick and Sweet J Biol Chem 80 52" (1998)

As a result the normal activity of the enzyme is decreased, and it produces an abnormal product which cannot be further metabolized and is therefore exceeded.

Valuable information has been gained from the study of those conjugated compounds which were the first historically to serve as metabolic tracers. Knoop as already stated, postulated the theory of β -oxidation from his observations of the end products of phenylaliphatic acid. Glycine, glutamine, ornithine, sulfune acid, and glucuronic acid were discovered as constituents of conjugated products long before it was recognized that these compounds participated actively in many normal physiological processes. Methylation of pyridine and nicotinic acid was noted decades before biochemists had any appreciation of the essential role of this reaction in metabolism.

The type of conjugation is determined by the active chemical groups in the molecule, but important modifications are brought about by second ary groups not directly concerned in the conjugation Marked species

differences in conjugation occur

The Carboxyl Group. The introduction of a carboxyl group into the benzene ring is one of the most effective means of reducing towict. Thus the addition of the COOH group to phenol produces saley lic acid, a non-toxic compound Toluene, which the body oxidizes to benzone acid, is less toxic than benzene, which yields on oxidation phenol polyphenols, and muconic acid. Benzone acid is the type compound in which a carboxyl group is attached to the benzene ring. In man nearly all of the compound is combined with gly cane to form hippuric acid. There is good evidence that the synthesis occurs primarily in the liver. The quantitative estimation of the excretion of hippuric acid following the administration of benzone acid is therefore a satisfactory test of liver function. With this method, the body's capacity to synthesize gly cine and to conjugate it with benzone acid are measured. The organism can utilize waste introgen for this synthesis, but the exact reactions in olved are not known.

Rather than postulate that the body possesses a mechanism specifically designed to synthesize hippure acid, it is more reasonable to suppose that benzoic acid has enough structural resemblance to a metabolite normally conjugated with giveine by a particular enzyme system to be acted upon by the same enzyme As a result benzoic acid is conjugated with glycine to form hippure acid, which is excreted because it cannot be further

setabolized. During the period that benzoic acid is being conjugated seenzyme is inhibited from carrying out its normal function. Evidence

of such a metaholic depression is the observation that the excretion of uric acid is strikingly decreased after the ingestion of henzoic acid.

Substitutions in the henzene ring of henzoic acid alter the pattern of conjugation. Any group in ortho position strikingly slows or inhihits the conjugation with glycine. This is well illustrated by the important drug, salicylic acid. Irrespective of the size of the dose, little of the compound is combined with glycine and excreted as salicyluric acid. Interestingly, when the hydroxy group is in para position, man excretes about one-half of the ingested compound unconjugated, and the other half combines with glycme to form p-hydroxyhippuric acid. p-Methoxyhenzoic acid, in marked contrast, is half conjugated with glucuronic acid and the remainder with glycine.

Phenylacetic acid differs sufficiently in structure from benzoic acid to require a different enzyme for its conjugation. This is demonstrated by the fact that when both kidneys are removed from a dog, the animal loses the power to synthesize hippuric acid, hut retains the ability to form phenylaceturic acid. It is of considerable interest that phenylacetic acid is the only compound known to he conjugated with glutamine This synthesis has heen observed only in man and in the chimpanzee. In the avian organism both henzoic and phenylacetic acid are conjugated with

ornithine.

Phenylacetic acid

The dog, in contrast to man and the rabbit, conjugates nearly 75 per cent of the ingested benzoic acid with glucuronic acid. The source of this compound remains unknown. Free glucuronie acid is poorly metabolized, and since it is not found in the unconjugated form, it seems unlikely that a

Giucuronic acid 1-Benzoyigiucuronic acid Benzole acid

direct union of glucuronic acid and benzoic acid occurs. The observation that the completely diabetic organism retains the ability to form conjugated glucuronides, apparently from the fraction which would be excreted as gluco-e, shows that glucogene amino acids can serve as a source of glucuronic acid. Whether the foreign compound is combined with gluco-e with subsequent oridation to glucuronic acid or whether the glucuronic acid is synthesized from three carbon compounds remains unsolved. The wide distribution of the enzyme β -glucuronidase in the body and the observation that various see hormones are combined and exerted with glucuronic acid clearly indicate that this compound occupies a prominent position in metabolism.

The Hydroxyl Radical. A compound such as phenol is conjugated with both glucuronic acid and sulfuric acid. The latter type of compound is called an ethereal sulfate. Indican, which is the common name for indoxyl-sulfuric acid, is lustorically important since it has been used as a measure of intestinal putrefaction. The organism apparently can combine sulfuric acid directly with a phenol Chloral hydrate (CCl₂Cli(OH)₂) a commonly employed sedative, is reduced in the body to trichloroethyl alcohol, which is conjugated with glucuronic acid.

The Aromatic Amino Group. Though aniline is extremely toxic, its acetyl derivative, acetanilide, is a relatively nontoxic drug. It would appear logical therefore to expect acetylation to be an important detoxication reaction. Actually there is little evidence that this is true Sulfanilar mide and its various derivatives are acetylated in the body. Their therapeutic effectiveness is thereby lost, but their toxicity does not appear to be significantly lowered. Interestingly, the vitamin p-aminohenzoic acid is acetylated and excreted. Although a etyleholine is an important physiological compound, acetylation of other hydroxy compounds, particularly nonmetabolites, has apparently not been observed.

suifonamide) The Pyridine Ring. Pyridine, quinoline, and many of their derivatives are methylated in the body. The transfer of a methyl group to nuclear nitrogen is apparently a common reaction. Nicotinic acid is methylated to form trigonelline. Since its antipellagra properties are thereby lost, the purpose of such a reaction is not clear.

Nicotinie acid

The Benzene Ring. When benzene or polycyclic hydrocarbons such as naphthalene and anthracene are ingested, their corresponding mercapturic acids can be isolated from the urine. In this conjugation, the sulfur of cysteme becomes directly combined with a nuclear carbon atom, and the amino group becomes acetylated. p-Bromophenylmercapturic acid formed from bromobenzene is the best known since it can be isolated fairly easily. Rather than regard this conjugation as a detoxication process, it seems more reasonable to suppose that it serves as a means for introducing hydrovy groups into the benzene ring, which is a step in the metabolism of the aromatic nucleus. This assumes that the body possesses an enzyme which splits the acctylated cysteine from the benzene ring by hydrolysis.

Bromobenzene The conjugation reactions should be regarded as important metabolic processes rather than biochemical oddities. The ease with which the body furnishes glycine, glucurome and, acetic acid, cysteine, and other conjugating components serves as evidence of their wide and important utilization in metabolism. Interestingly, the avian organism lacks the ability to form hippure acid, but conjugates aromatic acids with ornithme. This suggests the close relationship of some of the conjugating reactions with the Kirbe-Henselett urea cycle.

EXPERIMENTS

INDOLE AND SEATOLE

- Herter's β-Naphthoquinone Reaction. To 10 ml. of unknown solution add 2 drops of 2 per cent solution of β-naphthoquinone sodium monosulfonate and 2 ml. of 10 per cent NaOil. Let stand 15 minutes. Shake with 2 ml of chloroform. A pinkish-red color in the chloroform indicates indole. This is a very delicate test.
- Ehrlich's p-Dimethylaminobenzaldehyde Reaction. To 10 ml. of vn-known solution add 1 ml. of 5 per cent alcoholic solution of p-dimethyl-aminobenzaldehyde and 1 ml. of concentrated HCl. Indole gives a red color and akatole a blue color.

PHFNOLS, HYDROXY AROMATIC ACIDS, AND IMIDAZOLES

Conjugation with Glucuronic Acid and Glycine: Principle. The fate of an early (p-methoxybenzou acid) in the body illustrates the influence exerted by a relatively inter themical group such as the methoxy radical when introduced into the benacie ring of benzoue acid. In the human organism benzou acid is conjugated almost entirely with glycine to form hippure acid, whereas only about 50 per cent of p-methoxybenzoue acid is combined with glycine, the remainder is conjugated with glucurous acid. When glycine or gestam is given with the anisse acid, the amount of p-methoxyhippure acid exercted is greatly increased without, however, depressing the conjugation with fluvious acid.

by the naphthoresorcinol test. The remainder of the sample is acidlified with 1 ml. of concentrated hydrochloric acid and stirred to precipitate p-methoxy-hippuric acid. Inoculation with a small crystal of the compound will hasten precipitation. The crystalline product is filtered by suction, washed with a small amount of cold water, and dried. From the weight of the product plus the amount calculated that remained in solution (100 ml. of urine dissolve 0.24 g. of p-methoxyhippuric acid), the quantity of anisic acid which is conjugated with glycine can be calculated.

QUANTITATIVE BROMINATION

Method of Day and Taggard: Principle. Phenol, aniline, and many of their derivatives such as hydroxy- and sminoaromatic acids are readily brominated A quantitative replacement of one or more nuclear hydrogens by bromine occurs From the amount of bromine consumed, the quantity of aromatic compound is calculated

Procedure. 0.2 g. of p-methoxyhlppuric acid obtained in the preceding experiment and purified by recrystallization from hot water is neutralized with sodium hydroxide and diluted to 75 ml. The solution is quantitatively transferred to a 500-ml. glass-stoppered bottle, and 25 ml. of 0.2 N bromate solution (75 g. of KBr and 5.5 g. of KBr0, per liter) is added. The solution is acidfied with 5 ml. of concentrated hydrochloric acid and shaken for one minute. After 30 mlnutes the reaction bottle is cooled in ice or under the tap. The stopper is dislodged sufficiently to permit adding 5 ml. of 40 per cent RI solution. Care must be taken to prevent bromine vapor from escaping. The liberated lodine is titrated with 0 l N sodium thiosulfate, using starch solution as Indicator. The amount of bromine consumed is calculated from the difference between the number of ml. of thiosulfate required in the titration of the sample and the titration of the blank using 25 ml of 0 2 N bromate solution.

CALCULATION The bromine is supplied by the reaction

 $KBrO_1 + 5KBr + 6HCl = 3Br_2 + 3H_2O + 6KCl$

One molecule of p-methoxyhippuric acid reacts with one molecule of bromine

 $CH_4OC_6H_4COOH + Br_2 = CH_4OC_6H_4BrCOOH + HBr$

Each mi of 0 1 N throsulfate is equivalent to 0 01045 g of p-methox) hippuric acid

For the bromine equivalent of other amino and phenolic compounds the original articles of Day and Taggard and of Quick should be consulted

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⁴ Da) and Taggard Ind and Eng Chem. 20, 545 (1928), Quick J. Biol Chem., 97, 403 (1932).

21

Feces

General The faces include the residue remaining in the intestine after the digestion and absorption of food together with products of intestinal secretion epithelial debris, and bacterial growth and decomposition

They are composed of the following substances

1 Food residues 1e, those portions of the food which either escape of are meanable of digestion and absorption

2 The remains of the intestinal and digestive secretions not destroyed or realisorhed

3 Substances excreted into the intestinal tract, notably phosphates and other salts of calcium, iron, and other metals

4 The bacterial flora of the intestinal tract and their metabolic end products

5 Cellular elements to which may be added, under pathological conditions blood pus mucus serum and parasites

6 Abnormally, enteroliths, gallstones, and panereatic calculi

The amount of the feeal discharge varies with the individual and the diet. Upon an ordinary mixed diet various authorities claim that the dialy exercition by an adult malo will aggregate 110-170 g. with a sold content ranging between 25 and 45 g., the feeal discharge of such an in dividual upon a vegetable diet will be much greater and may even be a great as 350 g. and possess a solid content of 75 g. In the authors' experience the average daily output of most feces, calculated on the baus of data secured from the examination of over 1000 stools was about 100 g. The variation in the normal daily output is so great that this factor is of very little value for diagnostic purposes, except where the composition of

cruse the total amount of excrement to be markedly increased. An idea of the variation of the percentage of dry matter in the feces evacuated after the ingestion of different diets may be gathered from a consideration of the preceding table

Fecal Pigments The principal pigment of the feces is stercobilin which is chemically identical with the urobilin of urine and like it is formed by oxidation of a chromogen stercobilmogen (or irrobilmogen) This explains the darkening of stools upon exposure to air The primary precursor of these compounds is bilirubin which is reduced by intestinal hacteria to mesobilirubinogen, in the presence of a factor from bile this intermediate is transformed to urobilinogen or stereobilinogen according to the relations shown on p 413 A rapid dechne to neghgible amounts of urohihnogen in the feces bile and urine has been shown to follow the administration of aureomycin in therapeutic doses Neither bilirubin nor biliverdin occurs normally in the fecal discharge of adults although the former may be detected in the excrement of nursing infants If these pigments are found in the feces of adults they indicate an abnormally rapid transit through the large bowel preventing their transformation into stercobilinogen. A simple method has been suggested for the isola tion of stereobilin and urobilin from the feces in crystalline form The color of the fecal discharge is greatly influenced by the diet A mixed diet for instance produces stools which vary in color from light to dark brown an exclusive meat diet gives rise to a brownish black stool whereas the stool resulting from a milk diet is invariably light colored Drugs and certain pigmented foods such as cocoa beets the chlorophyllic vegetables and various varieties of berries each afford stools baying a characteristic color This is well illustrated by the occurrence of sellow stools following the administration of rhuharb senna or santonin and of red stools following prontosil treatment The green color of the colomel stool is generally beheved to be due to biliverdin The black stools following the administration of bismuth or iron drugs result from the formation of metallic sulfides or suboades In cases of bihary obstruction the absence of pigment and the presence of excess fat result in the formation of grayish white acholic stools Barnum meals used in roentgenographic diagnosis impart a clay white color to the fees

Odor Under normal conditions the odor of feces is due to skatole and mdole two substances formed in the course of putrefactive processes occurring within the intestine (see Chapter 20) Such compounds as methyl mercaptan hydrogen sulfide and similar substances resulting from bacterial putrefaction may also add to the disagreeable character of the odor The intensity of the odor depends to a large degree upon the character of the dict being very marked in stools from a meat dict much less marked in stools from a vegetable diet, and frequently hardly detectable in stools from a milk diet. Thus the stool of the infant is ordinarily nearly odorless and tends to be rancid rather than putrefac tive it is believed that any decided odor may generally be readily traced to some pathological source

Watson Lowry Shorov Hollmlead Kolan and Matte J Bud Chem 200 (97 (19.3)

Reaction. Experiments in which the actual hydrogen ion concentration of the feees was determined indicate that the normal reaction of the exerct a is slightly alkaline (pH 70 to 75) Pronounced dietary changes (e.g., low-protein diet, high protein diet, fasting, water-drinking with meals) produce at most only minor changes in the reaction of the fees The ingestion of large amounts of lactore may cause the production of an acid reaction

Consistency. The form and consistency of the stool is dependent in large measure upon the nature of the diet. Under normal conditions the consistency may vary from a thin, pasty discharge to a firmly formed stool. Stools which are exceedingly thin and watery ordinarily have a pathological significance. In general the feeces of the earnivorous animals

are of a firmer consistency than those of the herbix ora

The continued ingestion of a diet which is very thoroughly digested and absorbed is frequently accompanied by the formation of dry, hard, feed masses (exploid) Constipation generally results from the small bulk of the feces and its lack of moisture. At present the formation of sey bala is considered pathological, as an expression of spastic constipation. To counteract this tendency toward constipation the ingestion of agar agar psyllium seed or other vegetable gums is practiced. These are relatively indigestible and readily absorb water, thus forming a bulky feed mass which is sufficiently soft to permit easy evacuation. Mineral oil because of its lubricating and softening properties is much used as an aid in over coming constipation. Though it is insert and practically nonaborbable excessive amounts of mineral oil may interfere with the absorption of fat-soluble vitamins.

Separation It is frequently desirable for clinical or experimental pur poses to make an examination of the fecal output which constitutes the residual mass from a certain definite diet. Under such conditions it is customary to cause the person under observation to ingest at the begin ning and end of the period in question some substance sufficiently dif ferent in color and consistency from the surrounding feces to render comparatively easy the differentiation of the feces of that period from the feces of the immediately preceding and succeeding periods One of the most satisfactory methods of making this separation is by means of the ingestion of a gelatin capsule containing about 02 g of powdered char coal at the beginning and end of the period under observation This procedure eauses the appearance of two black zones of charcoal in the fecal mass A capsule containing earmine (0 3 g) may be used in a similar manner and forms two dark red zones Chromic oxide (Cr2O3) has also been suggested for this purpose * Some similar method for the separation of feces is usually practiced in connection with accurate nutrition of metabolism experiments conducted for the collection of useful data regarding the income and output of nitrogen and other elements

Macroscopical Examination Among the macroscopical constituents of the feess may be mentioned the following Intestinal parasites and their ova undigested food particles gallstones pathological products of

² Lloyd and Crampton J Audration 41 629 (19.0) Irwin and Crampton Ibid 43, 77 (1951)

the intestinal wall, enteroliths, intestinal sand, and objects which have heen accidentally swallowed

Microscopical Examination. The fecal constituents which at various times and under different conditions may be detected by the use of the microscope are as follows (1) Constituents derived from the food, such as muscle fibers, connective-tissue shreds, starch granules, and fat, (2) formed elements derived from the intestinal tract, such as epithelium, erythrocytes, and leukocytes, (3) mucus, (4) pus corpuscles, (5) parasites, and (6) bacteria In addition to the constituents named, the following

crystalline deposits oceasionally may be detected cholesterol, coprosterol, soaps, fatty acid, fat, bematordin, "triple phosphate," Charcot-Leyden crystals (see Fig. 104), and the oxalate, carbonate, phosphate, sulfate, and lactate of calcium

Fat in Feces. The amount and composition of fat excreted by way of the feces is largely independent of food fat but tends to approximate the composition of blood lipides Fecal fat is largely of endogenous origin, the secreted fat being partially reabsorbed, only to a minor degree does fecal fat come from epithelial debris and hacteria



Fig 104 Char COT-LEYDEN CRTS-

About one-third of the fat in normal feces is unsaponifiable. The coprosterol (also called coprostanol) of the feces is similar to cholesterol. and is formed by the reduction of the latter. It contains two more atoms of hydrogen than cholesterol and is thus a saturated alcohol C27H47OH A small amount of cholestanol, an isomer of coprosterol, is also found in feces It appears to be formed in the tissues and excreted in the bile Except for the D vitamins, ebolesterol is the only sterol absorbed from the intestine. The phytosterols of plants are not absorbed, being excreted in the feces Coprosterol responds to cholesterol color tests and has the same solubility, but possesses a lower melting point and crystallizes as fine needles instead of plates such as cholesterol forms

After the intravenous administration of cholesterol-4-C14 to normal rats it was found that from 10 to 14 per cent of the total C14 in hile and from 17 to 23 per cent of feeal C1 were recovered in the nonsaponifiable fraction This includes cholesterol, dihydrocholesterol, coprosterol, etc The fatty and fraction contained little if any C14 About 80 per cent of the fecal C14 and about 90 per cent of the bile C14 was present in the form of bile acids

The fat content of normal feces may vary from 5 to 25 per cent (on a dry basis) Excessive excretion of fat via feces is known as steatorrhea, a condition due to failure to absorb fat as in ententis, hepatic disorder, biliary obstruction, echae disease, or sprue Excreted fat is partly saponified and may entail loss of calcium, as soap, to the extent of producing pathological demineralization

Blood in Feces. The detection of minute quantities of blood in the feces-so-called occult blood-is an aid to the correct diagnosis of certain disorders. In these instances the hemorrhage is ordinarily so slight that

² Superatein Jay ko, Claukoff an i Dauben Proc Soc Fapil Biol Wed 81 720 (1952)

the identification by means of macroscopical characteristics as well as the microscopical identification through the detection of erythrocytes are both unsatisfactory in their results. Of the tests given for the detection of oecult blood the benzidine reaction is probably the most satisfactory. Since occult blood occurs with considerable regularity and frequency in gastrointestinal cancer and in gastrie and duodenal ulcer, its detection in the feces is of especial value as an aid to a correct diagnosis of these disorders. Certain precautions are essential, such as the establishment of a meat-free diet over a period of time before the specimen is collected. (Feces from a meat diet will give an occult blood reaction with some of the most delicate tests.) Bleeding from the howel such as is seen in hemorhoids, as well as the admixture of menstrual blood, is to be considered in the interpretation of the result. After the ingestion of 50 ml. of bursan blood 85 to 95 per cent of the hemoglobin appears as protohemin in the feces.

Bacteria in Feces. It has been quite clearly shown that the intestine of the newly born is sterile. However, this condition is quickly altered, and bacteria may be present in the feces before or after the first ingestion of food. There are three possible means of infecting the intestine; i.e., by way of the mouth or anus or through the blood. The infection by means of the blood seldom occurs except under pathological conditions; thus the usual sources are limited to the mouth and anus.

In infants with pronounced constipation, two-thirds of the dry substance of the stools has been found to consist of bacteria. In the stools of normal adults prohably about one-third of the dry substance is bacteria. The average exerction of dry bacteria in 24 hours for an adult is ahout 8 g. The output of fecal bacteria has been found to undergo a decrease under the influence of water-drinking with meals. There is also a decrease intestinal putrefaction, a fact which indicates that at least a part of the hacterial deficit is made up of putrefactive organisms. In some cases, more than 50 per cent of the total nitrogen of feces has been shown to be bacterial nitrogen.

Some of the more important organisms met with in the feces are the following: E. coli, B. lactis aerogenes, Cl. welchii, B. bifdus, and coreal forms. Of these the first three types mentioned are gas-forming organism. The production of gas by the fecal flora in dextrose bouillon is subject to great variations under pathological conditions; alterations in the diet of normal persons will also cause wide fluctuations. Data as to the production of gas are of considerable importance in a diagnostic way, although the exact cause of the variation is not yet established. It should be home in mind in this connection that gas volumes are frequently variable with the same individual. For this reason it is necessary in every instance to follow the gas production for a considerable period of time before drawing conclusions.

After aureomycin therapy, coliform organisms entirely disappear from the feces, while the clostridia diminish in quantity and may even disappear. No appreciable influence on the feeal flora has been shown to

⁴ Hughes: Brit. Med J., 2, 970 (1952).

follow the parenteral administration of penicillin and dihydrostreptomycin in combination

Enzymes in Feces. Various enzymes have been detected in the feces. The first one so demonstrated was pancreatic amy lase. The amy lase content of the feces has been considered to be an index of pancreatic activity. The excretion of this enzyme has been found to increase under the influence of water drinking with meals. Other enzymes which have been found in the feces under various conditions are trypsin, remnin, maltase, sucrase, lactase, nuclease, and house In an abnormally rapid transit of food through the intestinal tract, such as is seen in certain diarrheas, nearly all of these enzymes may be detected.

Vitamins in Feces. A variety of vitamins are present in normal feces. Their number and amount are subject to wide variation under pathological conditions. For example, the feces of enteritis patients have been found to be deficient in vitamins B₁ and B₆ (but not B₂) compared with feces of normal persons ⁶ Pennington ⁶ reports a heat-fable vitamin B₁₂ complex

m feces of the rat

Fecal Nitrogen. The nitrogen present in the feces exists principally in the form of (1) bacteria, (2) mabsorbed intestinal secretions and digestive juices, (3) epitbelial cells, (4) mucous material, (5) food residues, and (6) amino acids. In the early days of nutrition study the fecal nitrogen was believed to consist principally of food residues. We now know that such residues ordinarily make up but a small part of the total nitrogen of the stools of normal individuals who exercise normal mastication. When meat has been holted, bowever, from 0 5 to 16 g of macroscopical meat residues have been found in a single stool. The phrase "metabolic product nitrogen" has been used as a designation for all fecal nitrogen except that present as food residues and hacteria Bacteria cannot logically bo classed under metabolic nitrogen since they doubtless develop at the expense of food nitrogen as well as at the expense of that in the intestinal secretions In the accurate study of protein utilization? a correction should be made for metabolic nitrogen Data regarding the output of metabolic mtrogen may he secured by determining the fecal mtrogen excretion on a diet of proper energy value but containing no nitrogen However, to prevent tissue catabolism from exceeding its normal level, it is customary in studies of basal nitrogen metabolism to include a minimal percentage of adequate protein in the maintenance diet

Amino Acids in Feces. Fecal amino acids are derived from protein digestion as well as from intestinal organisms and secretions. It has been showns that everetion of free amino acids was more abundant in the feces of the breast-fed infant than in those of the artificially fed infant. The feces of the breast-fed infant showed alanine to be the predominant amino acid, whereas in the feces of the artificially fed baby value and lysine predominated. In experiments on white rats it has been shown that fecal excretion of lysine, value, histodire, and methioning is much higher on

Diaz et al Rev elin espai 44 233 (1952)
Lennington Biochem J 48 xxiii (1951)

Fee Protein Utilization in Clapter 33.
Ross Lancet 241, 190 (1951)

formalln or aicoholic solution of thymol (from which the aicohol may be evaporated by warming). Stools may be dried directly in these cans, If glass containers are preferred, pyres pots with glass lids may be used. To prevent loss of nitrogen sufficient sulfurle acid should be added to render the fecal mixture strongly acid. The contents may be dried on a steam bath in a bood and the residue weighed and powdered for sampling. If nitrogen or mineral analyses are to be conducted, the fresh stools may be mixed with 100 to 200 ml. of water and an equal volume of concentrated HySO, gradually added while stirring. The bomogenized suspension thus collected and preserved throughout the 24 hours may be diluted to volume and an aliquot taken for analysis.

- 2. Macroscopicol Examination. If the stool is watery pour it into a shallow dish and examine directly. If it is firm or pasty it should be treated with water and carefully stirred before the examination for macroscopical constituents is attempted. The macroscopical constituents may be collected very satisfactorily by means of a double layer of cheesecloth or a sieve of fine mesh.
- 3. Reaction. Thoroughly mix the feces and apply molst indicator papers to the surface. If the stool is hard it should be mixed with water before the reaction is taken. Examine the stool as soon after defecation as is convenient, since the reaction may change very rapidly The reaction of the normal stools of adult man is ordinarily neutral or faintly sikaline to litmus, but seldom acid. Infants' stools are generally acid in reaction. The glass electrode is useful in determining fecal pH since it is not affected by reducing substances commonly present, nor by the presence of solids.
- 4. Starch. If any Imperfectly cooked starch-containing food has been ingested, it will be possible to detect starch granules by a microscopical examination of the feces. If the granules are not detected by a microscopical examination, the feces should be piaced in an evaporating disb or casserole and boiled with water for a few minutes. Filter and test the filtrate by the fodine test in the usual way (see p. 84).
- 5. Blood. Undecomposed blood may be detected macroscopically II uncertain, look for crythrocytes under the microscope, and spectroscopically for the spectrum of ox) hemoglobin (see p. 492).

In ease the blood has been altered or is present in minute amount (occult blood) and cannot be detected by the means just mentioned, the following tests may be tried

BEADDER REACTION (a) Make a thin fecal suspension using about 5 ml. of their to remove the fat and discard this ether extract. Acidify the residue with acette acid and extract again with 5 ml of ether. Pour the acid ether extract into a small evaporating dish. Evaporate to dryness on a hot water bath (with flame turned out). Add a few drops of water, a drop of saturated solution of benzidine in glacial acetic acid and a drop of 3 per cent hy drogen peroxide. A blue or green color indicates the presence of blood.

(b) Supe Monreario. Take up a little of the solid stool on a match, a mear to nan object glass, and pour the reagent over it. If there is blood present the smear turns blue and there is no misleading green tint from fluid. Vlake the solution as follows; Add a knife-tip of benzidine to 2 ml. of glacial acetic acid, and and 20 drops of a 3 per cent solution of hydrogen percolde. By this dry technique there is no danger of soiling the fingers, and the test is more sensitive than the usual wet benzidine test. The smear of stool is either blue or it is not blue. The rapidity of the color change give some idea as to the proportion of blood in the stool; with much blood present the change to blue is instantaneous

(c) LEVIN-WATT MODIFICATION II Emulsify a small portion of feces in distilled water. Filter into a clean test tube, using a Whatman No. 5 or imore retentive filter paper. Treat 3 ml. of the filtrate with 8 drops of 50 pc cent acetic acid or glacial acetic acid, and mix. Add 8 drops of hy drogen perofilde (C P. 3 per cent) to the mixture and shake. Make a contact ring by carfully overlaying final mixture drop by drop with an alcoholic benzidine soft tion "Tilt the test tube in order to bring about slight mixing at the ring green color at the area of contact indicates a positive reaction. The intensit of color varies with the concentration of blood.

·42. 7

again added, and the cylinder shaken for five minutes and set aside. When the ether has nearly stratified, draw it off and wash as before. During the second washing, stratification will complete itself. Evaporate the ether until no trace of the alcohol which has been carried over with it remains. To the residue add 30 ml. of low-boiling petroleum ether (it should boil below 60° C), and allow to stand over night. Petroleum ether for this work should be frequently tested for a residue on evaporation. If a residue is left, the ether should be redistilled. Filter the petroleum ether solution of the fat, cutch the filtrate and washings in a tall, weighed, 100-ml. benker, evaporate off the solvent, dry at 90° C, desiccate, and weigh. After weighing, dissolve the contents of the beaker in 50 ml. of benzol, heat almost to the boiling point, add 2 drops of a 0.5 per cent solution of phenolphthalein, and titrate with a decinormal solution of sodium alcoholate.

CALCHATIONS The weight of total fat is obtained by subtracting the weight of the empty beaker from the weight of the beaker plus the dried fat. The weight of fatty acids (in terms of mg of stearin acid) is obtained by multiplying the number of ml of decinormal sodium alcoholate solution by the factor 28.4 The difference between the weight of total fat and the weight of fatty acids is the weight of neutral fat in the sample extracted

A separate determination without the addition of hydrochloric acid may be run upon the sample, for the purpose of determining the weight of neutral fat and free fatth acids. The difference between this weight and the weight of total fat is the weight of fatty acid present in the original sample in the form of soaps.

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¹¹ Erlenmeyer flashs of about 200 ml capacity may be used instead of beakers for the collection of the ether drawn from the eylinders. The ether may then be distilled and recovered The same procedure may be followed in removing the petroleum ether. **Rea Amendace**

22

Blood, Lymph, and Cerebrospinal Fluid

General. Blood serves as the principal transport system of the body, the heart furnishing the propulsive force. The most important quantitative functions of the blood are to bring oxygen and nutrients to the tissues and to earry away the waste products and deliver them to the excretory organs-the kidneys, lungs, biliary system of the liver, intestinal mucosa, and skin. Blood also plays an important role in coordinating the activities of various tissues through distribution of hormones, in maintaining the pH and the oxidation-reduction potential within narrow limits, in controlling temperature, osmotic pressure, and ionic concentration, in supplying defense against injection, and in guarding against hemorrhage.

The blood makes up about 5 to 7 per cent of the body weight in man, or approximately 2 to 3 liters per sq. m of body surface, being more nearly proportional to the latter than to weight. Lower relativo volumes of blood are observed in obese than in thun individuals. The volume of the circulating blood in a normal individual is maintained within rather narrow limits; any marked change in blood volume, as by sudden severe hemorrhage, etc., has serious consequences and may even result in death Blood volume may be measured by the addition to the blood of a known amount of a readily detectable and nondiffusible substance, such as certain dyes, hemoglobin derivatives, and radioactive compounds; the extent to which the substance is diluted by the blood is a measure of the total blood volume.

Composition of Blood. Since blood has numerous functions, its composition is necessarily complex. The general composition is summarized in the following table:

I. Cellular fraction Volume: 45 per cent

Constituent cells

- Erythrocytes (5,000,000 per emm.)
- 2 Leukocytes (6,000 per emm)
- 3 Platelets (250,000 per emm.)

II. Plasma fraction

Volume 55 per cent

- A. Nondiffusible constituents
 - 1. Albumins 2. Globulms

 - 3. Fibrinogen 4. Enzymes, lipides etc

B. Diffusible constituents

- 1 Catabolic products urea, creatimne, uric acid, etc 2 Anabolic constituents glucose, amino acids, creatine, etc
- 3 Electrolytes Na+, K+, Ca++, Mg++, Cl-, HCO₂, HPO₄, H-PO₄, etc

4 Hormones, vitamins, etc

The specific gravity of whole blood is about 1 060, that of plasma is approximately 1 026 Since the cells are heavier, they can be separated from plasma by centrifugation. The pH averages 7.4, the viscosity 1.7 to 2 (water = 1), the freezing point -0.56° C and the total osmotic pressure at 37° C is 7.6 atmospheres.

Blood Plasma and Serum When plasma is obtained without the ad dition of an anticorgulant, it is called nature Such a plasma, if kept from having contact with a wettible surface and covered with mineral oil to delay the escape of carbon dioude, retains fairly closely its in riso state and composition. For the usual chemical analysis, plasma obtained by the addition of sodium oxalate to prevent clotting is generally suitable and is commonly employed.

Scrum is the fluid obtained when blood clots. It is similar to plasma in composition but lacks fibringen and has a diminished concentration of other clotting factors such as prothrombin. Scrum is preferable to plasma for the determination of calcium albumin and globulin.

The plasma contains about 91 to 92 per cent of water and about 8 to 9 per cent of solid matter, of which about 7 per cent, or more than 85 per cent of the total solids, is protein

Plasma Proteins. The plasma proteins represent a complex mixture continuing a number of components which differ in properties and function. The major component proteins of plasma include (1) fibringen, (2) the various globulins, and (3) the albumins, (4) nucleoprotein and

(5) seromucoid are also present in limited amounts

TIBRINGEN The fibrinogen of normal human plasma is present in a concentration of about 0.19 to 0.33 g per 100 ml of plasma. The average in the Red Cross series was 0.28 g per 100 ml of plasma. The average found in lymph and ehyle as well as in certain evudates and transudates. Human blood fibrinogen is said to have a molecular weight of approunately 500 000, although research suggests that it may be as low as 350 000. The molecule is highly elongated, with a length estimated to be from 600 to 700 Angstrom units and with an avial ratio of about 20 to 1. The among acid composition is given in chapter 4.71brinogen is insoluble in salt-free water but is soluble in didute salt solutions. It is the most readily precipitable of all the common blood proteins by concentrated salt solutions, being precipitated upon half saturation with sodium chloride or 20 per cent saturation with ammonium sulfate. The coagulation temperature of fibrinogen is about 55° C in neutral solutions.

Tibrinogen is unique among the blood proteins in that it is readily converted into insoluble fibrin by the action of the enzyme thrombin Research has contributed to the elucidation of the mechanism of this reaction. Apparently when thrombin acts on fibrinogen, a fibrino-pertide is split off and the remaining molecule becomes activated. It polymer lizes to form fibrin, which separates as simple and compound fibers which interestingly exhibit cross-strictions with a constant periodicity. The fibrin fibers form a meshwork or reticulum which constitutes the framework of the clot of standing the clot shrinks, thereby expressing cles errum. This phenomenon, which is called clot retraction, depends on the presence of intact platelets which adhere to the fibrin strands and are some way responsible for condensing the fibrin mass.

Albumin and globulins The bulk of the plasma proteins consist of the albumin and globulin fractions. The albumin fraction ordinally preponderates comprising well over one-half of the total protein of normal human plasma, but this relation may be altered or even revired in disease. As a class the plasma albumins differ from the globulins in having a greater solubility, a lower molecular weight, and a more acid isoelectric point. Human serum albumin, for example, has a frolecular weight of 69,000 and an isoelectric point at pH 54, while the 7½ ulin of serum (see p. 463) has a molecular weight of 156,000 and in 1½ electric point at pH 65. Marked differences in amino acid composition between albumins and globulins have been established, as is cident from the data of the table on p. 122 in Chapter 4, but these difference while significant from an analytical point of view, as yet have not been related to differences in either structure or function.

The albumin fraction of the plasma is relatively homogeneous at dwell characterized, a number of serum albumins from virious sources have been obtained in crystalline form. Some of these crystilline proteins appear to contain carbohy drate as an integral portion of the molecule.

The globulin fraction on the other hand appears to consist of a variety of proteins of somewhat similar general characteristics but which by suitable means may be further fractionated into a number of components Although relatively homogeneous fractions of the plasma globulinhave been obtained, none of these proteins has as yet been crystallized From the point of view of solubility, two general types of globulin are recognized, englobulin and pseudoplobulin

I uzlobulm is a true globulm in that it is insoluble in salt-free water pseudoglobulm, while possessing the general properties of the globulms is soluble in salt-free water. This distinction between two types of globulms while useful, is by no means well defined. The extensive studies of Søren sen on this subject, already referred to in Chapter 5, have led him to postulate that the globulm fraction of sertim represents a loose combination of euglobulm and pseudoglobulm of the type EpP, in which F and P represent englobulm and pseudoglobulm complexes respectively combined in the relative proportions of p and q Fractionation by various means results in a shift in the proportions of E and P with the resultant formation of more soluble and less soluble complexes. Spirensen was unalle to prepare a sample of either englobulm or pseudoglobulm which we'll

1 Hawn and 1 orter J Prp Med 14 295 (191")

¹ Lorand Nature 147 992 (1931) Laskowski Rak witz and Sheraga J. Am. Chem. Soc., 74 28 (197)

entirely free of the other protein Cohn, McMeckin, et al., have likewise shown? that the englobulin fraction of scrum protein (i.e., globulin precipitated by dialysis) is considerably mereased in amount over that first obtained by ammonium sulfate fractionation if the precipitated protein is freed from the last traces of salt by electrodialysis and if the solubility is not influented by the presence of other proteins

Fractionation of the Plasma Proteins. The separation and characterization of the individual protein components of the plasma is of considerable importance. It facilitates study of the chemical nature and physiological function of each protein and of the significance of variation in the protein composition of the plasma in health and disease. It likewise kind to the possibility of the commercial preparation, from human or initial plasma, of purined preparations of the individual inoteins for laboratory, clinical, and industrial use

Methods for separating the plasma proteins from one another are based almost entirely upon (1) differences in physical properties, such as solubility in water, concentrated salt solutions, and other solvents; (2) rate of sedimentation in the ultracentrifuge, and (3) rate of electrophoretic migration. The various plasma proteins do not differ sufficiently in chemical composition or behavior to permit their separation at the present tune on a purely chemical basis. Immunological means of separation—1 et he use of precipitus (cf. Chapter 5)—specific for the individual protein, he we certain disadvantages and have been little used.

DIFFERENCES IN PRINCIL PROPERTIES The type of fractionation obtained by differences in solubility in water and in concentrated salt solutions is illustrated in the following table which has been adapted from the work of various investigators 'The salts listed in the table,

PRACTIONATION OF THE PLASMA PROTEINS BY CONCENTRATED SALT SOLUTIONS

Protein	Solubility in Dist H-O	Precipitated by		Approxi	
		Per Cent Saturation with NH ₄ Sulfate	G So- drum Sul fale per 100 ml, at 37° C	Normal Conc in g per 100 ml Plasma	Per Cent of Total Proteir
Fibrinogen Lugfobidin Pseudoglobidin I Pseudoglobidin II Albumin	- - + +	20 33 40 46 >50	13 5 17 5 21 5	0 3 0 2 1 3 0 5 5 2	4 3 17 7 69

ammonium sulfate and sodium sulfate, are those most commonly employed, sodium chloride, magnesium sulfate, and sodium or potassium phosphate my also be used under suitable conditions. It will be seen

¹ Cohn McMeekin Oncley Newell and Hughes J. Am Chem Soc. 62 3386 (1940). Howe J. Biol. Chem. 49 109 (1921). Gatman Moore Gatman McClellan and Kabri. J. Llin Intel. 20 763 (1941). and others.

that as the concentration of salt is progressively increased, the vanou fractions become insoluble and will precipitate from solution. Fibrinogen which is sometimes elassified as a globulin because of its solubility char acteristics, is the least soluble protein, followed by euglobulin, the pseudoglobulins, and albumin. It is usually considered that all the globulins of plasma are precipitated upon half-saturation with ammonius sulfate, or by a 22 per cent concentration of sodium sulfate at 37° C, and that the protein remaining in solution is albumin. This is the common analytical basis for separation of the albumin and globulin fractions of plasma prior to their analytical estimation.

The fractionation represented in the table above must be regarded as being quite arbitrary. The precipitation limits are not sharply defined, as might be inferred from the table, but rather represent arbitrarily established limits which in reality correspond to zones, between which there is no well-defined transition. The quantitative values given in the table are therefore characteristic of this particular type of fractionation only, and should not be expected to agree with values obtained by fractionation by other methods. Despite this limitation, the method has proved of value in many studies on the variation in plasma protein

fractions in health and disease.

Fractionation of the plasma proteins by various concentrations of aqueous alcohol is described by Cohn, Luetscher, et al. While alcohol denatures proteins readily at room temperature, denaturation does not occur at temperature ranges of 0° to -5° C., and satisfactory fractionation of the general nature of that olutamed by the use of concentrated salt solutions is obtained. The advantage of alcohol fractionation appears to lie chiefly in the case with which the fractionating agent may be removed from the protein, along with water, in the process of preparing stable dried preparations of the various fractions (lyophilization), thus rendering the method applicable to the large-scale preparation of such dry protein fractions for climeal and industrial use

By means of varying the temperature, the alcohol concentration, and the pH, Cohn and associate were able to reparate the plasma proteins into six fractions which they designated by Roman numerals Thus Fraction I, which contains most of the fibrinogen as well as a high concentration of the factor which is deficient in hemophila, is obtained at -3° C with 8 to 10 per cent alcohol at pH 72 To obtain the other fractions, the alcoholic concentration is progressuley increased and the pH is lowered Of particular clinical importance as Fraction V which contains nearly alf of the alhumin. None of the fractions, however, contain pure component Most of the reglobulm is detributed between Fractions II and III This globulm furnishes definite protection against the paralytic effect.

of pohomy clitis.

The low-temperature alcohol method was supposed to yield native serum proteins I infortunately, even though the albumins were still soluble at the isoelectric point they were denatured to the extent that they were poorly metabolized by the organism when injected. The low-

³ Cohn, Luctucher Oneley Assistrong and Davis J Am Chem Soc 42, 2396 (1940).

temperature alcohol method has therefore been supplanted by a procedure involving the use of specific adsorbants and fractional precipitation with zinc salts at pH 7.4. The Zn⁺⁺ ions are removed by means of a cation exchange resin. The following components have been identified in the fractions obtained in this method:

On Cation-Exchange	On Stroma of Red	Precipitate with	Filtrate from
Resin	Cells	Zn++ at pH 7.4	Zinc at pH 7.4
Protbrombin Protbrombin acceler- ator Leukocyte factor Lipoproteins	Isoagglutinins	Fibrinogen Immune globulins Other globulins \$\beta\$-Lipoproteins Cu-protein, etc.	Albumins Metal-combining globulin Glycoproteins Amylase Iodoprotein Esterase etc.

The serum albumin fraction so prepared is relatively heat-stable, does not contain any known virus contaminants, and is approximately as active osmotically as untreated human plasma.

Use of the Ultracentrifuge. Separation of the plasma proteins in the ultracentrifuge of Svedberg (see Chapter 1) has had somewhat limited applicability because of the superiority of other available methods, but has yielded valuable theoretical information. It has been found that bovine serum contains an albumin with a molecular weight of about 64,500 as estimated from the sedimentation data, together with several globulins. The two most abundant globulins have molecular weights of about 165,000.

ELECTROPHORFSIS. The fractionation of the plasma proteins by the use of electrophoresis (see Chapter 1) was initiated largely by the pioneer work of Tiselius. In the electrophoretic separation of the plasma proteins, advantage is taken of the differences in mobility of the various protein ions present under the influence of a potential gradient. The protein solution is first dialyzed against a suitable fulfer solution through eellophane tubing (Viskludier).



FIG. 105. TYPE OF CELL USED FOR ELEC-TROPHORESIS STUDIES OV PLASMA PROTEINS Courtesy, Klett Manufacturing Co., New York.

ing sausage casing) in order to equilibrate the two solutions with regard to their pH and conductivity. In an electrophoresis cell of the type shown in Fig. 105, one of the rectangular limbs and the bottom section are filled with the protein solution, while the other limb and the rest of the electrophoresis cell including the electrode vessels (not shown in the figure) are filled with the buffer solution serving as the outside fluid during dualysis. Sharp starting boundaries are produced by manipulating the cell sections. Reversible electrodes of the silver-silver elloride type, immersed in strong KCl solution, are connected with a suitable source of

high voltage direct current. Under the influence of the potential gradient bus produced (4 to 8 volts per em) the various species of protein ionare the migrate downward in one limb of the cell and upward in the other, in the direction toward the anode at alkaline pH values, at a rate which is a function of the surface-charge density, i.e., the number of free positive (or negative) charges per area unit of the surface of the protein molecules. By so moving, the individual protein ion species form more or less well-defined boundaries arranged in the order of their respective electrophoretic mobilities. However, it is to be noted that there is a considerable amount of overlapping of the various protein regions so that

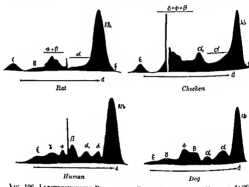


FIG. 106. I LECTROPHORETIC PATTERNS OF BLOOD PLASMA FROM VARIOUS ANIMAL SPECIES

Courtesy Deutsch and Goodloe J Bud Chem 161 1 (194o)

only the fastest and the slowest components in a multicomponent system may, as a rule, be obtained in pure form by means of the moving bond any technique. As the concentration of the individual proteins changes, the index of refraction of the medium likewise changes, and by suitable optical methods based upon this principle the migration of the violence protein fractions may be followed (Longsworth, Svensson)

The type of pattern obtained during electrophoresis of a mixture of components is illustrated by Fig. 106, which represents the electrophoretic patterns of vanous animal plasmas. The distance along the zaxis is a measure of the relative velocity of movement of the vanous ion spicies present, and the height of the peaks corresponds to the difference in refractive index between the moving boundary and the adjacent fluid, the area under each curve being proportional to the amount of material present moving with an average velocity represented by the

position of the peak along the x nus. Thus by this method it is possible not only to distinguish between ion species of different mobilities but also to estimate the relative amounts of each type of ion present

Tiselius demonstrated that normal plasma contained at least five electrophoretically distinguishable components, which were identifiable as albumin, fibringen, and three globulins, designated α -, β -, and γ -glob ulm respectively, the a globulin having the fastest velocity in the globuhin group and the y-globulin the slowest From electrophoretic data, it is clear that while albumin and fibringen are reasonably homogeneous—the high sharp peak corresponds to a migration of protein ions the majority of which have the same net charge and weight—the globulin fraction consists of a number of types of ions which may be classed roughly into three groups but obviously include ion species of quite varying mobilities within any one group Indeed, later workers have been able to show that the three globulin fractions of Tiselius may be resolved into further com ponents which have been designated α_1 - and α -globulin, β_1 - and β_2 -globu. lin, etc. In addition, stationary peaks, designated as δ and ϵ respectively are observed in the ascending and descending boundary diagrams (se-Fig 106) These have been shown to represent boundary anomalies rather than additional protein components. The fibringen boundary, usually designated by the symbol ϕ , is situated between the β - and y globulin boundaries Tollowing the example of Longsworth, most workers employ a diethylbarbitume acid-sodium diethylbarbiturate buffer system of pH 86 and 01 ionic strength which yields the highest degree of resolution of human serum or plasma diagrams

The type of quantitative distribution of the plasma proteins that is

obtained by electrophoresis is illustrated in the table below

DISTRIBUTION OF PROTEINS IN CITRATED PLASMA AS DETERMINED BY ELECTROPHORETIC ANALYSIS*

	Grams per 100 ml of Plasma		
Total protein	6 03	100	
Albumin	3 32	55	
α-Globulin	0 84	14	
β-Globulin	0.78	13	
γ-Globulin	0 66	11	
Fibrinogen	0 43	7	

^{*} Adapted from data of Cohn Oncley et al J Clin Invest 23 417 (1944)

It is to be noted that the three main globulin components are here present in approximately the same concentration, and furthermore that the ratio of albumin to globulin (the so-called "A/G ratio") is 1 15. This value is to be contrasted with the value of approximately 20 or higher which is accepted as normal for fractionation by sodium sulfate precipitation.

There is no simple relation between the fractions of the plasma proteins

obtained by salting out and those obtained by electrophoresis studies Thus Cohn, McMckin, et al (loc cit) have reported that at 34 per cen saturation with ammonium sulfate, the protein fraction precipitating consisted largely of y globulin, one-third of which was euglobulin-ie precipitated on dialysis At 40 per cent saturation with ammonium sulfat the fraction contained α- and β globulins as well as some γ-globulin Th. fraction likewise contained about one-third euglobulin Increasing the saturation to 50 per cent vielded a precipitate which was free from γ globulin but which contained both α - and β globulins, with only a fe per cent of the fraction insoluble in water Thus the two methods fractionation—salting out and electrophoresis—yield overlapping resuland are not directly comparable But it is to be noted that by suitable salting-out methods fractions can be prepared which are homogeneous electrophoretically and which correspond to the fractions obtained electrophoretically Thus Pillemer and Hutchinson have reported that at 0° C and pH 67 to 69, fractionation of human serum proteins with 42 5 per cent methanol yields A/G ratios which compare favorably with those obtained by electrophoretic analysis

Recently, methods for the separation of small amounts of proteins e.g., serum or plasma by electrophoresis in filter-paper strips (zone electrophoresis) have been developed by Durrum, Grassmann Ti-cliu and others. This technique, though not as strictly quantitative as the classical moving boundary method of electrophoresis, promises to have widespread application in clinical routine work. Fig. 57 (p. 164) shows the photograph of a developed paper strip after serum electrophoresis and the corresponding curve obtained by photoelectric densitometry of the stained protein bands. It should be pointed out that in zone or paper electrophoresis the individual bands visible after treating the fixed strip with suitable stains—e.g., Amdo Schwarz 10B—contain the actual protein components rather than represent their boundary against zones of different refractive index as in the moving boundary method of electrophoresis (compare with diagrams in Fig. 106).

At the present time it would appear that the electrophoretic method of analysis of the plasma proteins has its greatest value in studying the relative amounts of the various components present and the changewhich these components undergo during disease with a view to possible aid in diagnosis, but fractional precipitation is of value in the large-scale preparation of the individual components for industrial or clinical use the purity of the fractions being controlled by electrophoretic applications.

Origin of the Plasma Proteins. The liver is usually considered to be the site of formation of the plasma proteins although other parts of the body may also have a function in this connection. Fibrinogen synthesis appears to be dependent entirely upon the liver. When the liver is damaged experimentally his porsons the fibrinogen content of the blood falls returning to normal with liver regeneration and repair. If the blood of a normal animal is removed as much as possible and replaced by defibring

Pillemer and Hittel neon J Biol Chem 151 299 (1945)

ated blood, the fibringen level is restored to normal in a few hours, in the henatectomized animal this restoration does not occur

En idence relating the liver to the production of the other proteins of plasma is less definite. Liver atrophy is frequently found in experimental and chinical hypoproteinemia Since the albumin fraction appears to suffer most under these conditions, it would appear that the liver is associated with albumin synthesis, and furthermore that the factors controlling the synthesis of albumin are different from those for globulin, indeed. there is considerable evidence that parts of the body other than the hver are concerned with plasma globulin formation

The study of plasma protein production in experimental animals is facilitated by the use of the experimental technique known as plasmanheresis in which whole blood is removed from an animal, the blood cells separated from the plasma and resuspended in a suitable isotonic protem-free medium, and returned to the animal's circulation. Thus a plasma-protein deficit is produced which may be maintained at any desired point, and the extent to which the animal attempts to restore the plasma-protein level to normal may be used as an index of plasmaprotein formation under various experimental conditions. Using this technique, for example, it has been found that plasma protein production in the dog may be maintained at normal levels for many weeks by the intravenous administration of a mixture of purified amino acids as the sole source of dietary nitrogen, and that those amino acids in particular which have been found by Rose to be essential in the diet of the young growing rat (see Chapter 33) are likewise significant in the production of plasms protein in the dog

There is increasing evidence that the plasma proteins are in continuous metabolic equilibrium with other proteins and amino acids of the body This view, first postulated by Whipple and collaborators as the result of studies on protein regeneration in animals by the plasmapheresis technique, is supported by the work of Schoenbeimer et al., using the nitrogen isotope Nis When amino acids containing this isotope were fed to rats in nitrogen equilibrium, both the tissue proteins and the plasma proteins were found to incorporate the isotopic mtrogen rapidly and at about the same rate On discontinuing isotope administration, the isotope gradually disappeared from the plasma proteins, the estimated half life of the plasma-protein molecule being about two weeks. These and other experiments indicate that the plasma proteins in an animal are not static but are subject to continuous influence by dietary and metabolic factors

Functions of the Plasma Proteins. In addition to the specific physiological function of certain of the plasma proteins such as the role of fibringen and prothrombin in blood clotting and the role of the plasma globulus in immunological reactions, certain general functions of the plasma proteins are recognized These may be classified as nutritive and physicochemical

Schoenheimer Ratner Bittenberg and Heidelberger J Biol Chem 144 541 545 (1942)

NUTRITIVE FUNCTION The nutritive function of the plasma proteins has been definitely established largely through the work of Whipple and his associates. They have shown that the nitrogen (and protein) requirements of the fasting animal can be adequately supplied by the intravenous injection of plasma protein Since this injected protein disappears from the circulation in a short time it is presumably metabo lized Analyses of urmary nitrogen excretion after plasma protein injection indicate an increased catabolism of protein Whether this mean that the plasma proteins are directly utilized metabolically or are firs converted into tissue protein is not known, such a distinction would ap pear to be relatively unimportant in view of the existence of such dynamic equilibrium between tissue proteins and plasma proteins a has already been postulated

Physicochemical Functions The physicochemical functions of the plasma proteins are equally important Chief of these appear to be to aid in the maintenance of a normal blood volume and in the maintenance of a normal water content in the tissue fluids By virtue of their colloids dimensions the plasma proteins cannot normally diffuse through the blood capillary membranes into the relatively protein free tissue fluids. They thus exert an osmotic pressure which acts as a force tending to hold a certain volume of water within the blood This colloidal osmotic pressure or oncotic pressure, as it is sometimes called has a magnitude of about 25 mm Hg Although this is much less than 1 per cent of the total osmotic pressure of the plasma nevertheless it becomes the domi nant osmotic force in the blood capillaries since the other plasma con

stituents are freely diffusible across the capillary membranes

Of the various plasma proteins, albumin is by far the most significant in connection with osmotic pressure, being estimated to account for about 80 per cent of the total osmotic pressure of the plasma proteins Gram for gram, albumin is at least twice as effective osmoticalli as globulin, largely because of its relatively lower molecular weight. It has been shown that 1 g of plasma albumin is responsible for the retention of roughly 20 ml of water in the blood A loss of plasma protein and particularly of plasma albumin, therefore leads to a diminished blood volume, and this is presumably a major cause of the symptoms of hemor rhagie shock The administration of plasma protein especially plasma albumin concentrates has been shown to be of considerable value in both experimental and clinical shock

The colloidal osmotic pressure of the plasma proteins is opposed his the filtering force of the blood pressure which tends to drive water and dissolved substances across the capillary membranes into the tissue fluids At the arterial end of a blood capillary the blood pressure exceeds the colloidal osmotic pressure and fluid is forced from the blood into the tissues at the venous end the reverse action takes place fluid entering the blood from the tresues \ormally there is a balance between the two opposing forces and fluid distribution between the blood and tisues is normal If this balance is upset by such conditions as a low plasma protein content or an increased permeability of the capillary walls to protein, excessive amounts of water will accumulate in the tissues (edema). A plasma-protein deficit is therefore an important (although by no means the only) cause of edema. Edema due to plasma-protein deficit has been successfully treated by measures which restore the plasmaprotein level to normal.

A second physicochemical function of the plasma proteins is in connection with acid-base balance. Because the plasma proteins in solution at pH 7.4 are on the alkaline side of their isoelectric points, they exist to a certain extent as alkali salts. They thus act similarly to the alkali bicarbonate of the blood in furnishing base for the neutralization of acid, and indeed it has been shown that in the plasma the proteins are second

to bicarbonate in importance in this respect.

Other functions of the plasma proteins are also recognized. They aid in promoting the mobility of the corpuscles, since red cells sottle more rapidly in plasma than in protein-free isotonic solutions. The globulin fraction appears to be the carrier of the immune substances of the blood, and indeed the isolation and concentration of immune substances from human plasma is an important phase of fractionating the plasma proteins. The plasma proteins also combine with certain drugs and have thus been ascribed a vehicular function, although the precise significance of this property is not yet known.

The Erythrocytes. The crythrocytes or red cells make up about 45 per cent by volume of the blood. Variations from this value are frequently encountered; they are associated with changes in either the number of cells per unit volume of blood, or the size of the individual cells, or both. The relative cell volume of blood is determined with the bematocrit, a graduated tube which may be filled with whole blood and centrifuced. The volume of packed cells relative to the initial volume of blood is a measure of the cell volume. The top layer of packed cells is frequently almost colorless because of the predominance of leukocytes, which are not so heavy as the erythrocytes. The same phenomenon is noted when blood is allowed to clot without agitation; the lighter superficial portion of the clot is then known as the "buffy coat." The rate at which the erythrocytes settle, the so-called sedimentation rate, is applied clinically. Why the red cells settle out at different rates in the blood of various pathological conditions is not known. However, the rate is more rapid in tuberculosis, cancer, and acute inflammations. Newborn infants have a very slow rate, whereas normal men have a slower rate than normal women. The rapidity of the rate increases in menstruation and in normal pregnancy.

The crythrocytes are responsible for the opacity of blood. If blood is diluted with water this opacity disappears in a few moments, the fluid becomes translucent, and the blood is said to be hemolyzed or laked. On diluting blood with 0.9 per cent sodium chloride solution, however, no hemolysis occurs. The explanation for this lies in the osmotic behavior of the crythrocytes. Each cell may be considered as a miniature osmometer, the water content of the cell depending upon the osmotic pressure both of the cell contents and of the surrounding medium. If, for example, the cell is in a medium such as 0.9 per cent sodium chloride solution, which has the same osmotic pressure as the cell contents, the water

content of the cell will not change, nor will its size, and the medium is said to be isotonic with the cell If the osmotic pressure of the medium is greater than that of the cell contents water will be abstracted from the cell, it will decrease in size, and the solution is said to be hypertonic A solution which has a lower osmotic pressure than the cell contents F hypotonic, in such a solution, the cell will absorb water and swell the extent of swelling depending upon the degree of hypotonicity In a suff ciently hypotonic solution the swollen erythrocyte loses its ability to retain hemoglobin, and liemolysis results. The osmotic pressure at which hemolysis occurs is known clinically as the fragility point, and its deter mination has a certain diagnostic value

Other agencies besides osmotie-pressure differences will bring about hemolysis of crythrocytes These include alkali, ether, chloroform, sospabile salts, saponins, certain bacterial toxins, and snake venoms. In the instances hemolysis must be attributed to the actual modification of

destruction of the cell stroms

Human cry throcytes are nonnucleated biconcave disks with an average diameter of about $8 \mu (1 \mu = 0.001 \text{ mm})$ Nammalian erythrocytes var in size from species to species, ranging from 2 μ to about 9 μ in diameter In the blood of hirds, fishes, amphibians and reptiles the crythrocy tes are ordinarily more or less elliptical, biconvex, and nucleated

The number of crythrocytes present in human blood depends upon many factors, such as age sex, altitude, exercise, etc It is usually considered that the blood of a normal adult male contains 5,000 000 erythrocytes per cmm, for a normal adult female the count is 4,500 000 lnereased red-cell count is noted after blood transfusion, during residence at high altitudes, and after strenuous physical exercise, in the latter case a count of 7,040,000 has been observed. An increase is also noted in starva tion, after partaking of food, after cold or hot haths after massage, after partial asphyxia, and after fright as well as after the administration of certain drugs and accompanying certain diseases such as cholera, diarrhed dysentery, and yellow atrophy of the liver In polycythemia counts as high as 11,000 000 have been noted and values almost as high have been ohserved in cyanosis Experimentally, polycythemia may be produced in animals by the inclusion of cobalt salts in the diet A decrease in the number of crythrocytes occurs in the different forms of anemia, values as low as 500 000 per cmm or lower having been noted in pernicious anemia

Erythrocytes possess the property of grouping together in masses or "clumping" This action occurs normally on a microscopical scale and since the cell aggregates settle faster than the discrete cells clumping is a major factor in determining the sedimentation rate of the blood cells a characteristic of blood the measurement of which was shown by Fabraeus to have considerable climical value. If the clumping power of the cells is so enhanced as to produce macroscopically visible clumps the process is called agglutination Cells other than erythrocytes (e.g. bacteria) possess this property, when spoken of in connection with the blood the term hemagglutination is frequently used Observation of hemagglutina tion is the basis for the establishment of blood types, so essential in connection with blood transfusion, indeed, it is agglutination of crythrocytes which renders incompatible the bloods of donor and recipient. A substance which will bring about hemagglutination is said to contain hemagglutinins; if these are species specific they are known as isohemagglutinins. In human plasma the isohemagglutinins are associated largely with the β- and γ-globulin fractions. Hemagglutinins are abundant in the vegetable kingdom; for a demonstration of hemagglutination, see p. 486.

Composition of Red Cells. The red cells contain approximately 65 per cent water and 35 per cent solids. Of the solids, the red chromoprotein hemoglobin comprises about 32 of the 35 per cent; most of the remaining 3 per cent forms the stroma of the red cell, which consists largely of protein, phospholipide, and cholesterol. Inorganic ions in the red cell include potassium, chloride, bicarbonate, and phosphato; of these, potassium is present in largest amount, being comparable quantitatively, and in certain respects equivalent physiologically, to the sodium of the plasma. Organic constituents include various phosphate esters and certain enzymes, such as phosphatases and carbonic anhydrase; this latter is of particular significance in connection with the function of the red cells in the carriage of carbon dioxide by the blood (see Chapter 24). Certain diffusible substances such as glucose and urea are found equally distributed hetween cells and plasma when concentrations are expressed in terms of the water present; it is important to remember in such comparisons as this that on a percentage hasis the amount of water in the cells is much less than that in the plasma

Although erythrocytes contain oxidative enzymes (the "yellow enzyme" of Warburg was first isolated from horse red cells), the respiratory metaholism of these cells is very small. The major functions of the red cell appear explicable on a purely physicochemical hasis in terms of the role of hemoglobin and other substances present in the transport of

oxygen and carbon dioxide by the blood

Hemoglobin. Hemoglobin, the red coloring matter of the blood, is the most abundant protein in blood, being usually found to the extent of about 14 to 10 g. per 100 ml of whole blood. It is normally found entirely within the erythrocytes, from which it may be released by suitable hemolytic agents. It is a readily crystallizable conjugated protein, consisting of a colorless protein portion known as globin, which makes up about 96 per cent of the hemoglobin molecule, and a colored nonprotein portion or prosthetic group which has been shown to be an iron-containing compound belonging to the class of porphyrins. The chemical nature of globin has been discussed in Chapter 6.

The molecular weight of hemoglobin is approximately 67,000, concordant results being obtained by a variety of methods. The iron content of crystalline human hemoglobin is 0 340 per cent. Since this corresponds to a minimal molecular weight of 16,700 on the basis of one Fe atom per molecule, it has been concluded that the hemoglobin molecule contains four Fe atoms, and tho physicochemical behavior of hemoglobin is in

agreement with this conclusion.

It is now clearly established that several types of luman hemoglobin exist. In the prenatal period, the blood contains hemoglobin distinctly different from the adult type. It is called fetal hemoglobin and is recognized by its alkali resistance. Petal hemoglobin gradually disappears during the first year of life except in certain chronic anomina states in which it continues to be present in the blood. Siel le-cell anomia, a disease confined almost evolusively to the Negro is characterized by an abnormal hemoglobin and a peculiar behavior of the crythrocytes. The cells acquire a crescent shape when subjected to low oxygen tension. Siekle cell hemoglobin and normal hemoglobin have the same oxygen dissociation cure similar crystal forms and identical solubilities, but differ in their electrophoretic mobilities. This difference is sufficient to produce a serious disease which is transmitted by inheritance.

Hemoglobin is readily separated into its protein and prosthetic group components by treatment with acid, the globin usually being denature in the process and the iron containing portion being obtainable under the proper conditions in the form of an insoluble crystalline compound known as hemin. The formation of hemin crystals is frequently used as a test for blood because of their characteristic appearance and the case with which they may be obtained. Hemin has been synthesized and its structure is represented by the following formula.

It can be seen that the hemm molecule contains iron in the trivalent form and that the rest of the molecule consists essentially of four substituted methyl pyrrole rings linked together by —CH= bridges Such a molecule is known as a metalloporphyrin the metal in this case being iron Various isomers of the porphyrin molecule are known differing in the relative position of the substituting side chains the isomer found in hemin being called protoporphyrin To indicate the valence of the iron which is of importance it has been suggested that the terms ferror protoporphyrin and ferriprotoporphyrin be used Thus hemin is the chloride of ferriprotoporphyrin hemin them is also frequently used as equivalent to ferriprotoporphyrin hemin then being the chloride of heme. The older term "hemain bas been largely abandoned because of its indefinite connotation it is roughly equivalent to heme.

When hemin is treated with a suitable reducing agent the ferric iron is reduced to the ferrous state. The resulting ferroprotoporphyrin of reduced heme readily combines with undenstured globin to form a compound which is closely similar to if not identical with natural hemoglobin. Thus hemoglobin is a combination of globin with reduced heme

the iron in the hemoglobin molecule being in the ferrous state. The nature of the combination between the protein and the prosthetic group is not known.

Heme and reduced heme have the ability to combine with other nitrogen-containing compounds besides globin and denatured globin; such substances include other proteins, ammonia, cyanide, nicotine, pyridine, etc. These combinations are known in general as hemochromogens. Spectroscopic and other studies of the various natural and synthetic bemochromogens have thrown much light on our knowledge of the chemical behavior of heme and hemoglobin, and hemochromogen formation has been utilized for the qualitative detection and quantitative determination of hemoglobin. The occurrence of heme in nature as the prosthetic group of the enzymes catalase and perovidase, and in the evytochromes, has already been discussed (see Chapter 12).

Combination of Hemoglobin with Oxygen. A major physiological function of hemoglobin is hased upon its ability to react reversibly with

oxygen. This reaction may be written as follows:8

Hb $+ O_2 \rightleftharpoons \text{HbO}_2$ Hemoglobin Oxyhemoglobin

Under optimal conditions, 1 g of hemoglobin nill combine with 1 36 ml. of oxygen. The product of this reaction, oxyhemoglobin, is as well characterized a compound as hemoglobin, and is quite stable with respect to its oxygen content as long as there is sufficient oxygen present to prevent the reaction from going to the left Oxyhemoglobin may be crystallized, and a specife form of crystal is obtained from the blood of each individual animal species (see Figs. 107 and 108) Reichert and Brown studied oxyhemoglobin crystals prepared from the blood of more than 100 species of animals from the point of view of their crystallographic characteristics. Species differences are not confined to crystal form only; Barcroft and others have shown that oxyhemoglobin from various sources may differ in spectrescopic characteristics and in affinity for oxygen. Sinco hemin crystals are identical no matter from what species the blood is obtained, the species differences in oxyhemoglobins must presumably be related to differences in the globin portion of the molecule.

The reversibility of the reaction between hemoglobin and oxygen to form oxyhemoglobin implies that the relative amounts of hemoglobin and oxyhemoglobin present in blood will depend upon the concentration of oxygen present, which in turn is proportional to the oxygen tension (Henry's Law). The oxygen tension within the red cells is determined

largely by the oxygen tension of the plasma, since oxygen is freely diffusible across the red cell membrane

The relation between the degree of oxygenation of hemoglobin and the oxygen tension is usually expressed graphically in the form of a

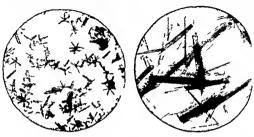


FIG. 107 OXYHEMOGLOBIA CRYSTAIS FIG. 108 OXYHEMOGLOBIA CRYSTAIS

THOM BLOOD OF THE RAT

These illustrat ons were reproduced from photom crographs furnuched by the late Prof. E. T. Pendert
of the Currently of Framylyins.

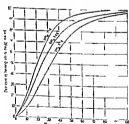


FIG 103 OXYGEN DISSOCIATION CURVE FOR HEMOCLOBIN IN BLOOD AT VARIOUS VALUES OF SERUM FII(pH)

Courtesy Van Slyke and Peters Quantitative Clinical Chemistry

curve the oxygen dissociation curve which is illustrated by Fig 109. The effect of such variables as pH and temperature on the ability of hemoglobin to combine with oxygen may be studied by noting their influence on the shape of the oxygen dissociation curve. From the curve it can be seen that when blood is in equilibrium with ordinary room air

(oxygen tension = ca. 150 mm Hg) or with the alveolar air of the lungs (oxygen tension = ca 90 to 100 mm Hg), practically all of the hemoglobin is oxygenated, i.e., the per cent saturation of the blood with oxygen is from 95 to 100 per cent. As the blood courses through the tissue capillaries, however, where there is a constant demand for oxygen, oxygen diffuses out of the plasma into the tissue fluids and the plasma oxygen

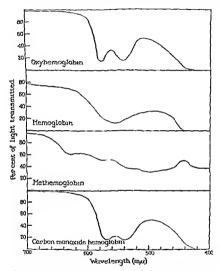


FIG 110 RELATION BETWEEN WAVELENGTH AND LIGHT ABSORPTION FOR HEMOGLOBIN AND OTHER BLOOD PROMENTS, ALL AT A CONCENTRATION OF 1 G, PER LITER AND AT A SOLUTION DEPTH OF 1 CM

tension falls to about 40 mm Hg. At this tension the oxyhemoglobin has yielded up about one-quarter of its combined oxygen by dissociation; thus normal venous blood contains about 75 per cent oxyhemoglobin and 25 per cent hemoglobin, or is 75 per cent saturated with oxygen Further lowering of the oxygen tension leads to increased dissociation of oxyhemoglobin; in the extreme condition of cyanosis only a minor portion of the pigment may be oxygenated. The sigmoid shape of the curve is considered to have some physiological significance. As the oxygen tension is lowered (i.e., as the demand for oxygen becomes greater), a given de

crease in tension produces a greater dissociation of oxyhemoglohin than is produced for the same decrease in tension at high tensions

The physiological function of hemoglobin is not confined to oxygen transport, it is of equal significance in the carriage of carbon dioxide by the blood. This function is discussed in detail in Chapter 24.

Other Reactions of Hemoglobin. In addition to its ability to read with oxygen, hemoglobin will react with a variety of other compound such as carbon monoxide, nitric oxide, hydrogen sulfide, ferricy and etc., the product in most cases being a colored compound which like hemoglobin and oxyhemoglobin, may usually be identified by its char



FIG. 111 CARRON MONOXIDE HEMO-GLOBIN CRYSTALS FEOM OX BLOOD Couriery Boor J Com Physiol. 13 207 (1930)

usually be identified by its classical acteristic absorption spectrum. The absorption spectrum. The absorption spectra of hemoglobin and some hemoglobin derivatives are shown in Fig. 110 as they are obtained quantitatively with the spectrophotometer The actual extent to which the light's absorbed is indicated by the data in Fig. 110.

The combination between hemoglobin and carbon monoxide to
form carbon monoxide hemoglobin
is of interest because of the constant presence of small amouats of
carbon monoxide in cits air Carbon monoxide hemoglobin is a
bright red pigment which may be
crystallized (see Fig 111) Carbon
monoxide has about 200 times the

affinity for hemoglobin that oxygen has, furthermore, the formation of carbon monoxide hemoglobin prevents hemoglobin from combining with oxygen, presumably because both gases compete for the same spot in the hemoglobin molecule. Carbon monoxide will in fact displace oxygen from oxyhemoglobin, as this reaction indicates

$HbO_2 + CO \rightleftharpoons HbCO + O_2$

This reaction ordinarily proceeds from left to right, it is however reversible if the concentration of oxygen is made high enough, and in this fact less the therapy for carbon monoxide poisoning. Carbon monoxide hemoglobin is not toxic in itself, death from carbon monoxide poisoning is presumably due to failure of sufficient oxygen to reach the tissues because of the decreased oxygen carrying capacity of the blood when a significant proportion (40 to 60 per cent or more) of the total hemoglobin is in the form of carbon monoxide hemoglobin. The anoxia of carbon monoxide poisoning is not due entirely to passive blocking of oxygen transport by hemoglobin, the oxyhemoglobin present in blood containing much carbon monoxide hemoglobin has been shown to be less efficient in releasing oxygen to the tissues under a given gradient of oxygen tension. The blood of city dwellers regularly contains over 1 per cent of the total because of the tissues under a given gradient of oxygen tension.

hemoglobin as carbon monovide hemoglobin, tobacco-smoking may in crease this to 5 per cent

When hemoglobin is treated with certain oxidizing agents either in tatro or in vito the substance methemoglobin is formed This is a brown pigment which differs from bemoglobin and oxyhemoglobin in that the iron is in the ferric form i e methemoglobin is a combination of heme or ferrinrotoporphyrin and globin Methemoglobin is usually detected by means of its characteristic absorption spectrum (see Fig. 110). It does not combine with either oxygen or carbon monoxide but does form a colored evanide derivative There is bttle or no methemoglobin in normal blood the appearance of a methemoglobinemia is noticed in certain diseases and after the administration of certain drugs among them sulfanilamide Methemoglohin is not toxic its presence in blood simply means a proportional reduction in the oxygen earrying capacity of the blood. The clinical induction of a moderately severe methemoglobinemia has been proposed as an aid in the treatment of cyanide poisoning since methemoglobin combines with evanide and may thus prevent the latter from re acting with enzymes in the tissues

Origin of Hemoglobin Hemoglobin is synthesized in the body from the ingredients of the diet iron a source of the porphyrin nucleus and amino acid precursors of the protein globin are of obvious importance in this connection It is known that inorganic iron of the dict cannot be converted into hemoglobin without the presence of catalytic traces of copper the way in which copper functions is not known 10 Studies on the origin of the porphyrin nucleus and on the significance of dietary amino acids in hemoglobili synthesis have yielded little definite information work with isotopes indicates that glycine and acetic acid may be pre cursors of the porphyrm ring (see also Chapter 33) In the adult animal the erythrocytes are formed in the bone marrow in the embryo the liver is an important source of red cell production and the possibility that in the adult the liver retuins a vital part in hemoglobin synthesis and erythrocyte formation cannot be overlooked. In the nutritional anemias there is usually a dietary deficiency leading to impaired hemoglobin formation when the deficiency is corrected the anemia disappears. In permicious anemia there appears to be lacking a factor which is necessary for the production of mature erythrocytes This factor vitamin Biz or cy mocobalamin (see p 1207), bas been isolated and found to contain 4 5 per cent of cobalt Interestingly both vitamin B1. and fohe acid correct the maturation defect in pernicious anemia but folic acid fulls to prevent the neurological complications which commonly occur. It is believed that the absorption of vitamin B12 is defective in permitious anomia and that this is due to the lack of a factor secreted by the stomach which is known as the intrinsic factor of Castle This would explain why vitamin B1 is exceedingly effective when given parenterally even in minute doses but relatively meffective when administered orally

The White Cells The white corpuscles (or lenkowites) of human blood differ structurally from the red corpuscles (or erythrowites) in many

Class Copper Metabolism Balti nore Johns Hopkins University Tress 1950 Marston Physical Revs 32 66 (1959)

particulars such as being larger in size, containing at least a single nucleus and possessing amedoid movement. They are typical animal cells and therefore contain the following substances which are customarily present in such cells Proteins, fats glycogen purines, enzymes, phosphatides cholesterol, morganic salts, and water Compound proteins make up the chief part of the protein quota of leukocytes, the nucleoproteins predominating Powerful proteolytic and gly colytic enzy mes are also present It is believed that there are two proteolytic enzymes in leukocytes, one active in alkaline solution and present in the polynuclear cells, and the other active in acid medium and present in mononuclear cells It is claimed that the granular leukocytes originate in the hone marrow, whereas the nongranular leukocytes (lymphocytes) have a lym phatic origin (lymph glands or lymphoid tissue), this matter of origin is uncertain The normal number of leukocytes in human blood varies between 5 000 and 10 000 per emm The ratio between the leukocy tes and erythrocytes is about 1 350 to 500

A leukocytosis is said to exist when the number of leukocytes is increased for any reason Leukocytoses may be divided into two general classes the physiological and the pathological Under the physiological form would be classed those leukocytoses accompanying pregnancy, partuntion, digestion, and excessive physical excreise as well as those due to mechanical and thermal influences Leukocytosis is also associated with such emotional states as fear, rage or apprehension The leukoeytoses spoken of as pathological are the infiammatory, infectious, posthemorrhagic, toxic, and experimental forms as well as the type which accompanies malignant disease

Chylomicrons Blood contains myrads of spherical particles, about 05 to 10 micron in diameter, which are highly refractive and show Brownian movement These were first recognized by Boyle in 1665, but were later described more fully by Muller who called them hemaconia or blood dust However, Gage showed them to be minute fat globules and curves of the chylomicron counts following the ingestion of fat have been studied in digestion experiments

It has long been known that hyperlipemia is common in diabetes and that the incidence of atherosclerosis is high in this disease. The relation slup of stcroids in the blood to arterial degeneration has aroused intense interest 11 Evidence is accumulating which suggests that steroid macromolecules may have an important causative role in atherosclerosis The original finding of Hahn12 (which has been repeatedly confirmed) that heparin has a striking effect on reducing postprandial hyperlipemia may nerhans be of both theoretical and clinical significance

Blood Coagulation When blood is withdrawn from a vein and put into a test tube it will clot solidly in less than 10 minutes. After about 30 minutes the coagulum begins to retract, with the expression of a clear serum These seemingly simple phenomena are the result of a complex scries of reactions The actual clot is due to the conversion of a soluble protein fibrinogen, to fibrin, which separates as fibrils forming a mesh

¹¹ Gofman Jones Lindgren Lyon Elhott and Strisower Circulation 2 161 (19-0) 12 Hal n Science 98 19 (1943)

work that incloses both the cellular and plasma elements of the blood Fibrinogen is changed to fibrin by the action of the enzyme thrombin For the formation of this clotting factor, at least five agents bave been shown to be essential prothrombin, labile factor, calcium, thromboplastinogen, and platelets

PROTHROUBIN Of the five basic thrombinogenic factors, protbrombin is the most clearly characterized and differentiated as a functional entity. It is a protein containing about 4 per cent earbohy drate. Its proceizyme nature is indicated by the fact that it becomes mactivated at 58 to 60° C. It is quantitatively removed from oxilated plusma by adsorbents such as tricalcium phosphate and barium sulfate and may be completely recovered from these adsorbents by elition with sodium citrate

Vitamin K is essential for the synthesis of prothrombin, but the amount needed is exceedingly small. An adult dog requires only about 05 \$\mu\$ go for natural vitamin K per kg of body weight daily to maintain a normal level of prothrombin. It is likely that the human requirements are about the same. The action of vitamin K can be satisfactorily explained by postulating that it functions as a prosthetic group combining with an apoenzyme (\overline{E}) to form the holoenzyme (\overline{E}), and that it is this enzyme which produces prothrombin. A deficiency of either apoenzyme or vitamin K results in a diminution of the holoenzyme and therefore in a decrease of the body's capacity to produce prothrombin. In the newborn a deficiency of vitamin K can easily occur due to inadequate intake. In the adult faulty absorption of vitamin K from the intestines is the most common cause for deficiency. Vitamin Ki, which is fat soluble, requires bile salts for its absorption. Any condition therefore, such as biliary obstruction in which bile does not enter the intestinal tract will bring about an inadequate absorption of vitamin K.

While the concept of an apoenzyme is still largely an bypotbesis the bypoprothrombinemia observed in certain liver diseases which is refractory to vitamin K is satisfactorily explained by the assumption that the apoenzyme is diminished Likewise, congenital hypoprothrombinemia, which is unaffected by vitamin K, can be attributed to a lack of the factor which with vitamin K forms the enzyme that produces prothrombin

Hypoprothrombinemia can be produced by a number of compounds which act as anti-vitamin K agents The best known of these is discoumard [3 3-methylenebis(4 hydroxycoumann)] which Link isolated from spoiled sweet-clover hay It may be postulated that discoumard depresses the production of prothrombin by replacing vitamin K in the enzyme system

Æ K according to the equation Æ K + D \rightleftharpoons Æ D + K, where D stands for dicoumarol The reaction is reversible and therefore the inhibitory action of dicoumarol can be counteracted by administering vitamin K Dicoumarol and several other vitamin K antagomists are employed thera peutically to produce hypoprothrombinemia as a means of lessening the tendency to thrombosis

Prothrombin is readily and accurately determined by means of the prothrombin time. In this procedure an excess of a standardized tissue extract such as acctone-dehydrated rabbit brain is added to recalcified plasma and the clotting time is determined By means of a curve, the prothrombin time can be expressed as per cent of normal clotting time Under the conditions of the test, the clotting or prothrombin time is a direct measure of the prothrombin concentration A second method known as the two-stage procedure consists in diluting plasma, after it has been defibrinated with a minute amount of thrombin, and converting all the prothrombin to thrombin by adding thromboplastin and calcium The thrombin is determined by the speed with which it coagulates a fixed amount of fibringen

Research indicates that only part of the prothrombin in human blood is in the active form, the remainder is present in a precursor atate to which the name prothrombinogen has been given. In the blood of the newborn bahy, prothrombinogen is lacking but the concentration of active prothrombin is the same as in adult blood 13 The prothrombin time which measures only free or active prothrombin and is uninfluenced by prothrombinogen, is therefore normal Since the two-stage method does not distinguish between free and inactive prothrombin but determines only total prothrombin, the concentration as determined by this method is only about one third of the adult level Since the newborn baby with a normal prothrombin time has no bleeding tendency, irrespective of the low prothrombin shown by the two-stage method it may be concluded that only active prothrombin participates in hemostasis and that there fore the prothrombin time is the more rehable test

Labile Factor When human oxalated plasma is stored at 4° C the prothromlan time becomes progressively prolonged. Since the addition of fresh plasma deprothrombinized by adsorption with Car(PO4); restores the prothrombin time of stored plasma to normal it must be concluded that the disappearance of a factor other than prothrombin must account for the loss of prothrombin activity during atorage. This agent has been named labile factor but is also known as Pactor V or Ac-alobulin (Ac accelerator) Like prothrombin it is a protein and is inactivated at 58 to 60° C Unlike prothrombin it is not adsorbed by Ca₂(PO₄)₂ nor is 1t5 concentration in the blood diminished by vitamin K deficiency or by dicoumarol The concentration of labile factor is relatively low in human blood as compared with rabbit blood which contains 50 times more yet a significant diminution of this agent rarely occurs clinically

CALCIUM Tike prothrombin and labile factor calcium reacts stoichiometrically in the formation of thrombin The minimal requirement for

¹¹ Quick Murat Hussey an I Burgess Surg Gynscol and Obst 95 671 (1952)

optimum clotting of human plasma is 0 0012 M. Since the level of calcium in the blood at which fatal tetany occurs is still entirely adequate for normal clotting, hypocalcemia as a cruse of bleeding is highly improbable

Prothrombin, labile factor, and calcium may for convenience be con sidered to constitute the prothrombin complex. The interrelation of these factors is stoichiometric, and since labile factor and calcium are normally in excess, prothrombin is the bmiting factor and therefore the determinant of the prothrombin time except in very rare pathological states

THROMBOPLASTINGEN Plasma contrins a protein, closely associated in physical properties with fibringen and globulin, which reacts with a platelet factor to form thromboplastin This agent, which has been named thromboplastinogen, is deficient in hemophilia. It is probable therefore that it is the same or closely related to the substance which some in vestigators call antihemophilic globulin

Curiously, no preformed thromboplastin is found in the blood Nevertheless, nearly all other tissues of the body contain this agent, and in some organs such as the lungs brain, and placenta it is present in rela-

tively high concentration

PLATELETS These cells contain a factor which reacts with thromboplastinogen to form thromboplastin The clotting agent occurs in the portion of the platelet called the granulomere. It is heat-stable and is probably a lipide with a high molecular weight. In addition to the participation of platelets in the formation of thrombin, they play an important role in clot retraction Intact platelets adhere to the shafts of fibrin and, probably as the result of morphological alterations, cause a twisting shortening and bending of fibrin strands that results in a condensation of the fibriu mass Platelets have a third function in hemostasis Lysis of these cells is accompanied by the appearance in the serum of a vasoconstrictor called serotonin or thrombotonin, which is chemically 5 hydroxy tryptamine It is probable that this compound evercises an important function in stopping the flow of blood in hemorrhage

THE CLOTTING REACTION The basic reason why circulating blood remains fluid is that it contains no active thromboplastin. For this substance to form, platelets must interact with thromboplastinogen but hefore even this can occur, thromboplastinogen must be activated by

means of thrombin The probable sequence of reactions is

(1) thromboplastinogen thromboplastinogen A (activated)

(2) thromboplastinogen A + platelet factor = thromboplastin

(3) thromboplastin + labile factor + calcium + prothrombin = thrombin

(4) fibringen thrombin fibrin

Thrombin has a dual function it clots fibringen and it activates thromboplastinogen Through the latter it initiates a chain reaction for as it activates thromboplistingen, more thrombin is formed, accelerating the cycle of reactions A break or injury of the vessel wall may liberate enough tissue thromboplistin to produce the initial thrombin that starts than 50 mg of protein per 100 ml., of which 80 per cent is albumia, the remander globulin

The osmotic pressure and the pH of the spiral fluid are the same as that of the plasms, whereas the gluco-e and calcium contents are roughly 50 per cent as great. The sodium chloride concentration is distinctly higher than that of plasma, while the bicarbonate concentration is about the same.

The analysis of the cercbrospinal fluid supplies valuable diagnostic information in the study of diseases involving the central nervous system In general, the quantitative methods developed for plasma can be applied with mmor modifications in the study of spinal fluid.

The colloulal gold test of Lange may also be used. In this test a colloids' gold solution is mixed with progressively increasing dilutions of cerebro spinal fluid. The orange-red color of Lange's solution is changed when cerebrospinal fluid from certain pathological conditions is tested

EXPERIMENTS ON BLOOD

IDENTIFICATION OF BLOOD

- 1. Microscopicol Examination. Place a drop of diluted blood on a microscopical siide and cover with a coverglass. Framine under a microscope.
- 2. Tests for Vorious Constituents. To 50 ml. of water in a large casserole. add 2 drops of acetic acid and hring to a boil. At the point of vigorous boiling. add 10 ml, of defibrinated blood slowly from a pipet, with stirring, Continue boiling and stirring for one minute after all the blood has been added. Then pour the mixture immediately onto a large folded filter which has been prepared beforehand. If the filtrate is not practically clear and colorless it should be discarded and the process repeated with more blood. Reserve the coagu lum for further use. Why is the coagulum colored dark brown? Evaporate the filtrate to about 25 ml., filtering off any precipitate which may form in the process. Make the following tests upon the filtrate:

(a) Bevenier's Test To 5 ml. of Benedict's solution add 5 drops of neutral lzed filtrate and boil one minute. Explain.

(b) CHLORIDES To a small amount of the fitrate in a test tube add a feet drops of nitric acid and a little silver nitrate. In the presence of chloride, white precipitate of silver chloride will form.

(c) Phosphares Test for phosphates by nitric acid and molybdate solution according to directions given on p. 213.

- (d) CALCIUM To 5 ml. of filtrate add 0.1 ml. of 4 per cent ammonium oxalate. Calcium oxalate precipitates as fine crystals. For crystalline form see p. 855 Y(e) Test for Inc. Incinerate a small portion of the coagulum obtained above, in a porcelain crucible. Cool, dissolve the residue in dliute hydrochloric acid, and test for iron by potassium ferrocyanide or ammonium thiocyanate. Which of the constituents of the blood contains the iron?
 - (I) CRISTALLIZATION OF SORIUM CHLORIDE Place the remainder of the filtrate in a watch glass and evaporate it on a water bath. Examine the crystals under

(a) TEIGIMAN'S METHOO Place a very small drop of blood on a microscopical slide, add a small drop of water, and stir to lake the blood Add a fraction of a drop of dilute (0 9 per cent) sodium chloride or potassium chloride solution.

and carefully evaporate to drvness over a low flame. Put a coverglass in place, run underneath it a drop of glacial acetic acid, and warm genth until the formation of gas hubbles is noted. Add another drop of glacial acetic acid, cool the preparation, examine under the microscope, and compare the crystals with those shown in Fig. 113.

(b) Alppe's Mericon Spread a small drop of blood on a slide in the form of a film and evaporate to dryness over a low flame Now add 2 drops of a solution containing 0 1 g each of potassium chloride, lodide, and bromide in 100 ml of glacial acetic acid Place a coverglass in position and heat gently over a low flame



Fig 112 Sodium Chlorice

until gas bubbles form and the solution boils Run I to 2 drops of the rengent underneath the coverglass and examine under a microscope Compare the crystals with those shown in Fig. 113



the clotting reaction. The prompt removal of thrombin is therefore necessary to safeguard against uncontrolled and massive intravascular clotting Experimental evidence has been obtained indicating that the physiological antithrombin is fibrin

Owing to the ability of fibrin to adsorb thrombin with avidity and because of the tremendous surface it presents to the serum dispersed in the reticulum of the clot the removal of thrombin by fibrin is exceedingly efficient and the chain reaction is effectively held in abevance Since the intimate contact between scrum and fibrin surface is lost when clot re traction occurs, it is very likely that this process may play a significant role in intravascular clotting

There is increasing evidence that the clotting reactions in addition to their function in hemostasis may perhaps participate in immunological and other defense mechanisms. It is probably not accidental that throm hoplastin is widely distributed in various tissues and that the daily turnover of prothrombin is nearly equal to the amount in circulation As the physiological significance of blood clotting becomes better understood a number of factors directly or indirectly associated with these reactions may be better evaluated Among such factors is the antithrombin of the serum This agent, which occurs in the albumin fraction neutralizes large quantities of thrombin, but its action is so slow that it does not interfere with the conversion of fibringen to fibrin and therefore has no true anticoagulant action

Fibrinolysin is an enzyme which dissolves fibrin. It is present in the blood as a 73 mogen, profibrinolysin Interestingly certain bacteria produce products which can activate profibrinolysin Such activators are

streptokinase and staphylokinase

Another factor which markedly influences the clotting of blood 15 heparin a mucopolysaccharide containing mucoitin sulfuric acid Heparin is present in the mast cells of Fhrlich as polychromatic-staining granules The amount occurring in the blood is extremely small, but in anaphylactic shock its concentration may become sufficiently high to make the blood incoagulable Heparin itself has little anticoagulant action but it forms a powerful antithrombin with a cofactor which is present in the albumin fraction of serum

Medicolegal Tests for Blood The analysis of a specimen for the presence of blood may be of utmost significance Often it is even more im portant to determine whether the blood is of human origin To establish the presence of blood a portion of the material under investigation should be extracted with 0.9 per cent sodium chloride solution and examined under the microscope II the material contains relatively fresh blood erythrocytes may be found and identified. An aqueous extract should be examined spectroscopically or better extracted with acid alcohol and examined with the spectroscope for absorption bands of acid hematin The preparation and identification of bemin cry stals is a satisfactory way to detect blood Even old blood will readily yield these dark brown or chocolate crystals, provided the specimen has not been exposed to a high temperature or to direct sunlight for a long period If the amount of blood is very small detection by the guarac test or the benzidine reaction is of great practical usefulness. These tests are exceedingly delicate and reliable, but it is essential that the reagents he properly prepared and tested on control solutions of highly diluted blood.

After the presence of blood is established by these tests the final step is its identification as to species origin. This can be done only by immunological procedures of which the precipitin test is the most sintisfactory. This test is founded on the observation that when serum of one animal is injected into an animal of a different species, the latter will develop in its serum an agent which will react with the proteins of the foreign serum and cause the formation of a precipitate. The precipitin test is highly delicate. It will detect blood in high dilution. It is likewise very specific, since overlapping occurs only in very closely related species.

Lymph. Lymph may be considered the middleman in the transactions hetween blood and tissues. It is the medium by which the nutritive material and oxygen transported by the blood for the tissues are brought into intimate contact with those tissues and thus utilized. In the further fulfillment of its function, the lymph hears from the tissues water, salts, and the products of the activity and catabolism of the tissues, and passes these into the blood. Lymph therefore evercises the function of a gohetween for blood and tissues. It hathes every active tissue of the animal hody, and is believed to have its origin partly in the blood and partly in the tissues.

In chemical characteristics, lymph resembles blood plasma. In fact, it has been termed "blood without its red corpuseles." Lymph from the thoracic duct of a fasting animal or from a large lymphatic vessel of a well-nourished animal is of a variable color (colorless, jellowish, or slightly reddish) and alkaline in reaction to litmus. It contains fibrinogen prothrombin, and leukocytes, and coagulates slowly, the clot heing less firm and bulky than the blood clot. Serum illumin and serum globulin are both present in lymph, the allumin predominating in a ratio of about 3 or 4.1. The principal inorganic salts are sodium salts (cbloride and hicarhonate), the phospbutes of potassium, calcium, magnesium, and iron are present in smaller inmount.

Substances which stimulate the flow of lymph are termed lymphagogues Such substances as sugar, urea, certain salts (especially sodium chloride), peptone, egg albumin, extracts of dog's liver and intestine,

crab muscles, and blood leeches are included in this class

In a fasting animal, the lymph coming from the intestine is a clear, transparent fluid possessing the characteristics already outlined. After a meal containing fat has been ingested, this intestinal lymph is white or milky. This is termed chyle, and is essentially lymph possessing an abnormally high (5 to 15 per cent) content of emulsified fat. This chyle is absorbed by the lacted of the intestine and transported to the lower portion of the thoracic duct. Apart from the fat content, the composition of lymph and chyle are similar.

Cerebrospinal Fluid. On tapping the spinal can'd a water clear fluid is obtained which is essentially a plasma filtrate formed by the choroid pleans. This fluid fills the two lateral and the third and fourth ventroles the subarachnoid, and the spinal canal Spiral fluid usually contains less

4 Gualac Test This methol is sensitive and valuable for detecting blood Care must be taken not to use too concentrated a solution of guriac since a voluminous precipitation of resinous material may easily obscure the blue color. Substances other than blood such as milk pus and saliva may give a positive test, but after boiling for 15 to 20 seconds they no longer yield a blue color with guaiac, whereas blood similarly subjected to boiling still yields a positive test

The test follows By means of a pinet drop a solution of gualac in glacial acetic acid (strength about 1 60) into the solution under examination14 until a turbidity is observed, and add hydrogen peroxide, drop by drop, until a blue color is obtained

5 Benzidine Reaction This test is one of the most delicate for detecting blood but care must be taken to use a good grade of benzidine, since the sensitivity is greatly influenced by the purity of the reagent. The test depends on the action of hemoglobin which catalytically decomposes hydrogen peroxide, thereby liberating oxygen which oxidizes benziline to a blue or green derivative. Since an excess of hydrogen peroxide interferes with the reaction it is essential that the details of the procedure be followed scrupulously It is particularly important that the peroxide be added last. The benzi dine solution is unstable, especially when exposed to light. It should therefore be prepared fresh daily and kept in a brown bottle or a dark place. The test is particularly valuable to detect occult blood in the feces 15

The test is performed as follows To 3 ml of a saturated solution of benzi dine in glacial acetic acid" add 2 mi of the solution to be tested and 1 mi of 3 per cent hydrogen peroxide A positive test is indicated by a blue or green color The following modification is a much more delicate and reliable test

Confirmatory Test Make 10 mi of the solution acid with acetic acid and extract by shaking with 5 ml of ether The acid breaks up the hemoglobin to globin and heme and the latter is extracted by the ether Pour off the ether into a small evaporating dish Put on a hot water bath (with the flame turned out) Fyaporate to dryness To the residue add a few drops of water, a drop of saturated solution of benzildine in glacial acetic acid, and a drop of 3 per cent hydrogen peroxide A blue or green color indicates blood

Lyle, Curtman, and Marshal Modification Into a perfectly clean dry test tube introduce 1 4 ml of benzidine solution, " add 8 2 ml of water or glacial acetic acid, then 1 ml of the fluid to be tested and finally 6 4 ml of 3 per cent hydrogen peroxide Note the appearance of a blue color, which reaches its maximum in 5 to 6 minutes

6 Hemochromogen Test Add 2 or 3 drops of Takayama's solution18 to 2 small piece of suspected material on a silde Cover with a coverglass Examine

¹⁴ Alkaline solutions should be made slightly acid with acetic acid since the blue end reaction is very sensitive to alkali. This is particularly necessary if an alcoholic solution of guarae is used

¹⁴ Hugles Brit Med J 2 970 (1952)

¹⁴ Glacial acetic acid is preferable but if it is not available alcohol acidified with acetic acid may be used

W Benz dine solution may be prepared as follows: Place 4.33 ml of glacial acetic acid in a small Erlemmeyer flask warm to 50° and add 0.5 g of benzidine. Heat the flask for eight to the minutes in water at 50° To the resultant solution add 10 ml of distilled water. This solution may be kept for several days without deterioration

18 A mixture of 3 ml of 10 per cent NaOH 3 ml of pyridine 3 ml of a saturated solution

of glucose and 7 ml of water The solution works rapidly in the cold if at least 24 hours old With a fresh solution warming or more time is necessary. It keeps for from one to two months.

under the microscope. Salmon pink crystals should appear in 1 to 6 minutes. At the same time the color changes through green-brown and dark red to pink, indicating the formation of a hemocbromogen and confirming the test. The crystals have a shallow rhomboid form (see Fig. 114).

This method is simpler to use than the hemin test and may be used to confirm the latter in doubtful cases. It is not always given by old blood stains (over six months old) and hence does not replace the bemin test. The glucose may act as a reducing agent as well as by decreasing the solubility of the



FIG. 114. HEMOCHEOMOGEN CRYSTALS

Prepared and photographed by Dr. Arthur G. Cole
University of Illinois, College of Medicine

hemochromogen. Schumm has cast some doubt on the specificity of this test for blood,

7. Immunological Determination of Blood Species. The serum proteins give immunological reactions, such as the precupitur reaction, which are specific not only for the individual proteins of the serum but also for the species of animal from which the serum is obtained. These immunological reactions are therefore used in medicolegal investigations to determine the species to which a particular blood sample or stain belongs. The species specificity of these reactions is demonstrated in the following experiments on the precipitur reactions of dog, beef, and human serums. Antiserums against human blood react with the serums of some of the higher apes, indicating a close relationship between the various species. Usually, however, even these reactions show quantitative differences which are sufficient to differentiate between human and other bloods.

(a) PREPAIATION OF IMMUNE SERUMS (ANTISERUMS). Prepare approximately 1 per cent solutions of dog, beef, and human serum proteins by diluting 15 ml. of the clear serums to 100 ml. with physiological (0.9 per cent) salt solution. Immunize rabbits against each of these solutions (antigens) by injection of the diluted serums into the marginal ear vein. Make the first injection with 1 ml. of antigen and follow with injections of 2, 4, 6, and 8 ml. portions at latervals of three or four days. On the fourth day after the last injection, collect 1 ml. of blood and test the serum for precipitin content by the method

described helow If the precipitin is present in sufficiently high titer, bleed the rabbit from the heart, allowing the blood to coagulate Transfer the clear serum to sterile vials closed with rubber stoppers. Add a few drops of chiloroform to each vial as a preservative, and keep vials in a refrigerator when not in use

An alternative method for preparing the indiscrums is as follows. Twenty-five mi of serum is diluted with 80 ml of distilled water, and to this solution 90 ml of 10 per cent potassium alum is added. The pil is adjusted to the isoelectric point specific to the alum protein complex (human serum, pil 6.5) with 5N NaOil. The precipitate is washed twice with 200 ml saline containing 1 10,000 merthiolate, and then is auspended in 100 ml of saline (1 10,000 merthiolate).

The immunizing dose is 10 ml, of the suspension, which is injected intramuscularly—5 ml into each hind leg of a rabbit, After 15 to 20 days the rabbit is taken off food for 24 hours and then bled The serum obtained is tested for specificity and sensitivity. For a positive result a precipitin at a dilution of a 1 100 antigen should be obtained within five minutes. As a control, an antigen of a species to which the rabbit is not sensitized should show no precipitin in a dilution of 1 50

(b) Tituation of Antherous Prepare the following dilutions of the antisea (diluted serum or protein solution to be tested) using physiological salina addition 1 100, 11,000, 110,000, 150,000 Transfer 1ml of each dilution to a separate small test tube which must be acrupulously clean and dry. With a capillary pipet carefully introduce 0.2 ml of the immune serum at the bottom of each tube so as to stratify the antigen solution above the serum With a 1 1,000 dilution, a white ring or cloud will form at the interface in a few minutes if the test is positive

For control, place in one test tube 1 ml of a i 50 dilution of an antigen against which the rabbit was not immunized, and add 0 1 ml of the antiserum as in the test above For a second control, transfer 1 ml of the 1 1,000 dilution of the antigen to a test tube and add 0 i ml of normal rabbit serum No precipitation should form in either tube

(c) IMMUNOLOGICAL EXAMINATION OF BLOOD STAINS Extract the stain with 1 or 2 ml of physiological salt solution. Files and

2 ml of physiological salt solution Filter and use the filtrate for making up dilutions of antigen and for precipitin reactions, as described above
8 Hemagglutination The common garden bean such as the scarlet runner,

contains a protein substance which exhibits the interesting property of causing clumping or agglutination of red blood corpuseles.

Dilute defibrinated bloods are also as the scarlet runner, and a substance which is a substance which is the standard of the substance with the sub

Dilute defibrinated bloods ten times with physiological sodium chloride solution (0 9 per cent) and place 1 ml in each of three small test tubes

Grind 3 beans to a fine meal in a coffee mill or with mortar and pestle and extract for a few minutes with 9 per cent sodium chloride solution Filer and add 0.5 ml (about 2 to 3 drops) of the filtered extract to the first of the blood tubes, 0.01 ml to the second, and 0.05 ml of 0.9 per cent sodium chloride to the right.

Invert each tube to mix the contents thoroughly, and note the rapid agglu tination and precipitation of the blood corpuscies in the first tube, a less

¹⁸ Habbit's blood is especially desirable and may be obtained for the purpose by bleeding from a small cut on the animal's ear and defibringing

¹⁸ The scarlet runner is a familiar variety purel another in every seed store. It occurs in two varieties the white and the red. Been as protein constituent of the eastor been also possesses pronounced agglutinating properties. Recause of its poisonous nature it is however, not suitable for use in class experiments.

rapid agglutination in the second, while the third or control tube remains unaltered in half an hour the corpuscles in the first tube often are packed solid and one is able to pour off perfectly clear serum

- If the remainder of the bean extract is bolled for a few minutes, the coagulum filtered out and 0.05 ml of the filtrate added to the control tube, no agglutination occurs, indicating that the hemagglutinin has been destroyed or removed by the boiling
- 9 Red Ceil Fragility When red ceils are placed in a hypotonic saline solution, water present into the ceil until the osmotic pressure within and without are equalized. This brings whout a swelling of the red ceil which if sufficiently great causes the ceil membrane to rupture, thereby resulting in hemolysis. Normal erythrocytes remain intact in as low as 0.44 per cent saline whereas defective red cells, present in certain types of disease such as hereditary benolytic jaundice, show beginning hemolysis at this concentration and complete hemolysis at 0.34 per cent.

Procedure Prepare a 1 per cent solution of sodium chloride and from this make a series of dilutions ranging from 0 30 to 0 58 per cent by placing 0 60 ml in the first test tube and 0 64 ml in the second, increasing each succes sive tube by 0 04 ml until the fifteenth, to which I 16 ml is added Add distilled water to each tube to bring the volume to 2 ml To each tube add one drop of blood obtained by renipuncture 'Mx and allow to stand at room temperature for 2 hours Record as beginning hemolysis the tube showing a slight tinge of red in the supernatant, and as complete hemolysis, the tube in which all cells have disappeared

10 Crystollization of Oxyliemoglobin (Reichert's Metliod) Add to 5 ml of the blood of the dog, horse, guinea pig, or rat, before or after laking or defibrinating, from 1 to 5 per cent of ammonium oxalate in substance Place a drop of this oxalated blood on a silde and examine under the microscope The crystals of oxylemoglobin will be seen to form at once near the margin of the drop, and in a few minutes the entire drop may be a solid mass of crystals Compare the crystals with those shown in Figs 107 and 108, p. 472

In some species (eg, the rat) oxyhemoglobia tends to crystallize out of blood very rapidly Merely pressing a small drop of blood between a covergiass and a microscope slide will result in a mass of oxy hemoglobin crystals

- 11 Preparation of Hemoglobin (Method of Marshall and Welker) Draw blood into a flask and defibrinate by shaking with glass beads. Strain through cheesecloth. Centrifuge Wash corpuscles three or four times with 0.9 per cent NaCl. Add ether a few drops at a time with thorough mixing until a clear solution is obtained if the solution is viscid add a little water and then an equal volume of aluminum hydroxide cream (see Appendix). Wit thoroughly and filter Cool to 0°C and add absolute alcohol (also cooled to 0° to make the alcohol percentage 20 to 30 per cent. Let stand at a few degrees below 0°C. Wash the crystals by decantation with 25 per cent alcohol at 0°C. Dry in a desicator over sulfuric acid
 - 12 Demonstrations of Hemolysis and Osmotic Pressure
- (a) Hemolesis (Laking Bloop) Note the opacity of ordinary defibrinated blood Place a few mi of this blood in a test tube and add water, a little at a time, until the blood is rendered transparent Hemolysis has taken place How does the water act in causing this transparency? Pramine a drop of

hemolyzed blood under the microscope. How does its microscopical appearance differ from that of unaltered blood? What other agents may be used to bring about hemolysis?

(b) OSMOTIC PRESENTE Place a few mi of blood in each of three test tubes Hemolyze the blood in the first tube according to directions given in (a), above Add an equal volume of isotonic (0 9 per cent) sodium chloride to the

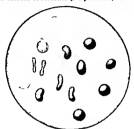


FIG. 115 DEFECT OF WATER ON PRYTHEOCYTES.

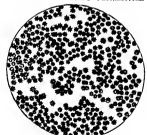


FIG 116 CRENATED ERTTHEOCYTES

blood in the second tube, and an equal volume of 10 per cent sodium chloride to the blood in the third tube 'lix thoroughly by shaking, and after a few moments examine a drop from each of the three tubes under the microscope (see Fig. 115 and 116) What do you find and what is your explanation from the standpoint of somotic pressure?

PLASMA AND SERUM

Plasma is the noncellular fraction of the blood. It is conveniently obtained by centrifugation. If no anticoagulant is used, the plasma can be

designated as native To obtain such plasma, it is best to collect the blood with a silicone-coated syringe and needle²¹ and to transfer it to a tube similarly coated. Usually sodium citrate or oxalate is mixed with the blood to make it incoagulable. It is customary to name the plasma according to the anticoagulant employed, as for instance, oxalated, citrated, or heparin plasma

Serum is the fluid obtained after blood has clotted. It has essentially the same composition as plasma except that it lacks fibringen and has a diminished concentration of various clotting factors.

- Preporotion of Oxalated Plasma. Collect 9 ml. of blood by venipuncture
 and mix immediately with 1 ml. of 0.1 M sodium oxalate¹¹ in a test tube or
 centrifuge tube. Centrifuge at 2500 r.p.m. for 5 minutes. Remove the plasma
 with a pipet. By attaching the rubber bulb of a medicine dropper to the pipet,
 the plasma can readily be drawn off.
- 2. Preparation of Serum. Place approximately 5 ml. of freshly drawn blood in a small test tube. Put the tube in a water bath at 37° C. for 30 minutes and then centrifuge at 250 r.p.m. for 10 minutes.
- 3. Seporotion of Fibrinogen, Globulin, and Albumin. Add 0.5 ml. of half-saturated ammonlum sulfate³³ to 0.5 ml. of oxalated plasma. Note the amount of precipitated fibrinogen formed. Repeat adding 0.5 ml. of saturated sodium chloride³⁴ to 0.5 ml. of oxalated plasma. Compare the amounts of fibrinogen obtained by the two salting-out methods.

Repeat the experiment, adding half-saturated ammonium sulfate and saturated sodium chloride to equal volumes of serum. Why do the solutions remain clear?

To 0.5 ml. of serum add 0.5 ml. saturated ammonium sulfate and mix. A precipitate of globulin is formed. Remove the salted-out protein either by centrifugation or fittration. To the clear filtrate, add small quantities of powdered ammonium sulfate and note the formation of a precipitate, which is albumin.

Outline a scheme of separating fibrinogen, globulin, and albumin.

Test samples of the precipitated fibringen, globuiln, and albumin with the bluret, Millon's, and Hopkins-Cole tests.

4. Heat Caugulation of Plasma Proteins. Transfer 0.5 ml. of plasma and 0.5 ml. of serum to separate tubes and place in a water bath at 60°C. Examine after 10 minutes. Serum remains clear because it does not contain fibringen. Gradually increase the temperature of the bath and note when heat coagulation appears in the serum.

²¹ Coaing glass with silicone Dilute one volume of Silicone (meth)1-chlorosilane-Dri-Film N 9937, General Electric) with 3 volumes of toluene Thoroughly clean the glassware and dry Cover the surface to be coated with the silicone solution Pour off the excess and drain for 30 minutes Rinse in warm running water and dry. To obtain a thorough costing, polish the surface with a njon cloth and recoat several times

n Sodium ozalate, 0.1 M. Dissolve 1.34 g sodium oxalate (cp) in 100 ml of distilled water.

11 Saturated ammonium sulfate Cover 90 g of ammonium sulfate (cp) with 100 ml

of distilled water. Vix and allow to stand for 21 hours. The supernatant liquid is saturated. Dilute with distilled water to obtain any desired saturation.

²⁴ Saturated sodium chloride. Cover 40 g with 100 ml. of distilled water, mlx, and allow to stand 24 hours

BLOOD COAGULATION

1 Clotting Time of Recalcified Plasma On adding calcium to citrated or oxalated plasma, clotting will occur. The probable over all equation to express the reaction is

thromboplastin + labele factor + release + prothrombin = thrombin

How these factors interact is not known, but there is strong evidence that the composite reaction is essentially storchiometric. The circumlaphistic does not exist preformed in the blood but is the product of the interaction of a platelet constituent and thromboplastinogen. The relatively slow and small production of thromboplystin is the limiting factor in the reaction and is therefore the determinant of the elotting time of recalefied plasma. If blood is oxalated in a test tube conted with insternal groung a nonwettable surface such as paraffin or silicone the platelets are preserved and can be removed by high centrifugation. I lasma thus obtained clots much more slowly on recalification.

Procedure Transfer 0 1 ml of oxalated plasma to a small test tube and place in a water bath at 37°C Blow 0 2 ml 0 01 M calcium chlorider into the plasma to attain instantaneous miting, and time the clotting exactly with a stop watch Examine the tube every 15 seconds hy gently tilting to detect the exact moment the inciplent clot appears Normal human oxalated plasma collected in glass clots in 90 seconds to 120 seconds after recalcification

To determine the effect of the removal of platelets on the clotting time-collect 10 ml of blood by venlpuncture Transfer 0.5 ml of 0.1 N sodium oxalate to a glass test tube and the same quantity to a test tube coated with parafin Add 4.5 ml of blood to each tube and mix by inverting the tube after covering it with wax paper. Chill the blood by placing it is n an ice bath for 10 minutes. Centrifuge the parafin coated test tube at 2,000 r p m for 15 mlnutes, and the glass test tube at the same speed but only for 3 minutes. Determine the clotting time of both plasmas on recalcification by the method as outlined.

2 Prothrombin Time The relative slowness with which oxalated plasma clots after recalcification is due to the small amount of available thromboplastin. By adding an excess quantity of a potent standardized thromboplastin reagents such as is readily prepared from rabbit brain and a fixed amount of calcium, the clotting time can be made a quantitative measure of it be prothrombin concentration, since the calcium and thromboplastin are constants in the reaction.

Procedure Transfer 0 1 ml of exalated plasma to a small test tube, add 0 t ml of thromboplastin reagent, and place in a water bath kept at 37° C Blow 0 1 ml of 001 ll catclium chtoride measured in a short pipet into the

the Calcium chlorule 0 01 M. Dissolve 0 11 g. in 100 ml. of distilled water is Preparation of Thromb oplastin Reagent. Remove the brain of a freely killed rabbit and

clear of all visible blood weeks Triturate the material in a mortar with 20 ml acctions and the material in a mortar with 20 ml acctions and the material becomes flax power of the aperial except a continuous properties of the material becomes flax power of the aperial acctions and again triturate. Hereat until the material is granular and nonadhenive. Filter by the continuous properties of the properties

Wir 200 mg, of the sectors dehydrated fram with 5 ml 0.85 per cent sodium chloride and incubate at 50° C for 20 muites. Wir occasionally by blowing through with a pipel Mace in a water bath 37° C and allow to settle sufficiently to permit drawing the solution into a pipel, and 37° C and allow the settle sufficiently to permit drawing the solution into a pipel.

mixture. Click the stop watch at the moment the calcium chloride is added and time accurately the appearance of the clot. This is hest done by leaving the test tube in the water bath until shortly before clotting occurs, then holding the tube towards a good light, and tilting gently to note the inciplent formation of fibrin. Normal fresh oxalated human plasma has a prothrombin time of 11½ to 12½ seconds. When the plasma is mixed with saline in varying proportions, the following values are obtained.

Dilution of Plasma	Concentration of Prothrombin	Prothrombin Time
	per cent	seconds
0	100	12
4 to 5	80	13
3 to 5	60	14
23∕2 to 5	50 40 30	15 17 19½ 24 to 26
2 to 5		
11/2 to 5		
1 to 5	20	
1∕2 to 5	10	37 to 42

By means of a serological pipet graduated in 0.1 ml. prepare the above dilutions and determine the prothrombin time of each mixture.

3. Preparation of Thrombin (Eagle). Dilute 10 ml. of fresh citrated or oxalated plasma with 100 ml. of cold distilled water. Chill the mixture to about 5° C. in lee water and then hubble carbon doxide gas through the solution for 5 to 10 minutes. Centrifuge off the precipitate and discard the supernatant fluid. Dissolve the residue in 10 ml. of 0.85 per cent sodium chloride solution and adjust the resulting solution to approximately pH 7.0 by the addition of sodium bicarbonate. Add to this final solution one-twentieth its volume of 0 1 M calcium chloride solution (1.1 per cent). Warm to 37° C. for a few minutes, insert a glass rod into the clot which forms and wrap the clot around the rod until all entrained fluid has been pressed out. Discard the clot, filter the remaining fluid, and store in the refrigerator. If kept cold, this thrombin solution is stable for about a week. Dried preparations of thrombin in sealed containers, which are stable indefinitely, may now be obtained from wholesale drug supply houses.

GLOTTING TYMES. When 0 1 ml. of this thrombin solution is added to 0.2 ml. of oxalated plasma, coagulation occurs in about 3 seconds. Prepare the following dilutions of thrombin with distilled water-1:2, 1:5, 1:10, 1:20 and 1:40. Determine the thrombin time of each dilution as follows Transfer 0.2 ml of oxalated plasma to a test tube, place in a water bath at 37° C. Blow 0.1 ml, of the thrombin solution into the plasma and accurately determine the clotting time with a stop watch. To prepare a curve, plot the clotting times against the dilution of thrombin, taking the undiluted thrombin preparation as 100

4. Preparation of Fibrin. Allow blood to flow directly from the animal into a ressel and rapidly whip it by means of a bundle of twige, a mass of strong cords, or a specially constructed beater. If a pure fibrin is desired it is not hest to attempt to manipulate a large volume of blood at one time. After the fibrin has been collected it should be freed from any adhering blood clots.

and washed in water to remove further traces of blood. The pure product should be very light in color, it may be preserved under giycerol, dilute alcohol, or chlorolorm water.

(a) Solubility Try the solubility of small shreds of freshly prepared fibrin in

water, dllute acid, and alkali

(b) PROTEIN COLOR TEXTS Test a portion of fibrin by Millon's test, the Hopkins-Cole test, and the bluret reaction. What amino acids have you shown to be present in fibrin?

SPECTROSCOPIC EXAMINATION OF BLOOD

As indicated in the text on p. 471, the spectroscope is a useful tool in the study of the blood pigments because of the differences in light absorption

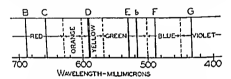


FIG. 117 DIACRAMMATIC REFERENTATION OF SOLAR SPECTRUM.

Shown are approximate extent of colored zones and location of principal Fraunhofer lines (B, C D E b F, G)

which these pigments show, and which are evident upon spectroscopic examination Fig. 117 shows a diagrammatic representation of the visible spectrum as seen with a spectroscope, including the approximate location and width of the various colored zones of the spectrum. If a solution containing a colored substance is placed between the spectroscope and the source of light, one or more dark zones or bands are seen in the spectrum, corresponding to the absorption of light of certain specific way elengths by the colored substance. It is frequently possible to distinguish between various pigments by the location and intensity of these absorption bands



FIG 118 DIRECT VISION SPECTROSCOPE

Since the ordinary hand spectroscope is not equipped with a wavelength scale, the absorption bands may be located by reference to certain of the more prominent Fraunhofer lines of the sun's spectrum These are dark lines, readily visible in the spectrum of sunlight, which correspond to the presence of certain elements in the vapors surrounding the sun As shown in Fig. 117, the most prominent lines and their approximate wavelengths are as follows B, 687m C 656m D 589m E, 527m D, 517m F,



FIG 119 ANOULAR VISION SPECTROMETER

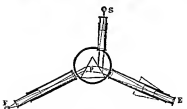


Fig 120 Diagram of Anoular Vision Spectroscope (Long)

The white light F, enters the collimator tube through a narrow slit and passes to the prism P, which has the power of refracting and dispersing the light. The rays then pass to the double convex lens of the ocular tube and are deflected to the eyepiece E. The dotted lines show the magnified virtual image which is formed. The third tube contains a scale whose image is reflected into the ocular and shown with the spectrum. Between the light F, and the collimator slit is placed a cell to hold the solution undergoing examination.

Either the direct vision spectroscope (Fig. 118) or the angular-visiospectroscope (Figs. 119 and 120) may be used in making the spectroscopic examination of the blood. For a complete description of these instruments the student is referred to any standard textbook of physics or to the catalogs of the manufacturers.

- Oxyhemoglobin. Examine dilute (1:50) defibrinated blood spectroscopically. Note the broad absorption band between D and E. Continue the dilution until this single broad band gives place to two narrow bands, the one nearer the D line being the narrower. These are the typical absorption bands of oxyhemoglobin obtained from dilute solutions of blood. Now dilute the blood very freely and note that the bands eradually become narrower and, if the dilution is sufficiently great, finally disappear entirely.
 - 2. Hemoglobin (So-called Reduced Hemoglobin). To blood which has been dlluted sufficiently to show well-defined oxyhemoglobin absorption bands, add a small amount of Stokes's reagent.27 The blood immediately changes in color from a bright red to violet-red. The oxyhemoglobin has been reduced through the action of Stokes's reagent and reduced hemoelobin has been formed, by the removal of the oxygen from the oxyhemoglobin. Examine this solution spectroscopically. Note that in place of the two absorption bands of oxyhemoglohin we now have a single broad band lying almost entirely between D and E. This is the typical spectrum of hemoglobin. If the solution showing this spectrum be shaken in the air for a few moments, it will again assume the bright red color of oxyhemoglobin and show the characteristic spectrum of that plement.
 - 3. Corbon Monoxide Hemoglobin. The preparation of this pigment may be easily accomplished by passing ordinary illuminating gas" through defibrinated ox blood. Blood thus treated assumes a brighter tint (carmine) than that imparted by oxyhemoglobin. Examine the carbon monoxide hemoglobin solution spectroscopically. Observe that the spectrum of this substance resembles the spectrum of oxyhemoglobin in showing two absorption bands between D and E. The bands of carbon monoxide hemoglobin, however, are somewhat nearer the violet end of the spectrum. Add some Stokes's reagent to the solution and again examine spectroscopically. Note that the position and intensity of the absorption bands remain unaitered.

The following are delicate chemical tests for the detection of carbon monoxide hemoglobin.

ALKALI TEST Mix two drops of the suspected blood on a porcelain plate with an equal volume of 25 per cent NaOII. A reddish color remains in the presence of carbon monoride hemoglobin. Treat two drops of normal blood in the same way. A brownish color is obtained.

TANNIN TEST Divide the blood to be tested into two portions and dilute each with 4 volumes of distilled water Place the diluted blood mixtures in two small flasks or large test tubes and add 20 drops of a 10 per cent solution of potassium ferricyanide." Allow both solutions to stand for a few minutes. then stopper the vessels and shake one vigorously for 10 to 15 minutes, occasionally removing the stopper to permit air to enter the vessel 20 Add 5 to 10 drops of ammonium sulfide (yellow) and 10 ml, of a 10 per cent solution of tannin to each flask. The contents of the shaken flask will soon exhibit the formation of a dirty olive-green precipitate, whereas the flask which was not shaken and which, therefore, still contains carbon monoxide hemoglobin will exhibit a bright red precipitate, characteristic of carbon monoxide hemoglobin. This test is more delicate than the spectroscopic test and serves to

¹⁷ See Appendix

[&]quot; The so-called water gas with which ordinary illuminating gas is diluted contains usually as much as 20 per cent of carbon monoxide (CO)

This transforms the oxyhemoglolan into methemoglolan which does not combine with

This is done to dissipate any carbon monoride present.

detect the presence of as low a content as 5 per cent of carbon monoxide hemoglobin.

DILUTION TEST In very dilute solution oxyhemoglobin appears y ellowishred, while carbon monoxide hemoglobin under the same conditions appears pinkish- or bluish-red. Dilute a drop of normal blood with water, and dilute in parallel fashion a drop of blood containing carbon monoxide hemoglobin with water, until by comparison a difference in tint is noted. This is said to be as satisfactory a test as any for routine purposes.

QUANTITATIVE DETERMINATION OF CARBON MONOXIDE 11 If possible collect blood from an arm vein in an oxalated tube. Otherwise wrap a finger of the subject with a rubber band. Prick deeply with a blood lancet. Draw 0.1 ml, of blood into a pipet and discharge into a test tube containing 0.9 ml, of distilled water, drawing back once or twice to insure removal of adhering blood, Mix immediately but not too vigorously. Add 1 ml, of freshly prepared pyrogallictannic acid solution, and mix by inverting twice. After 15 minutes compare with standards prepared as follows: Lake I volume of oxalated human blood free from carbon monoxide with 9 volumes of distilled water. Saturate half of the laked blood with CO by rotating in a flask or separatory funnel filled with illuminating gas for 15 to 20 minutes, avoiding violent agitation. Mix proportional parts of the treated and untreated blood solutions to represent 0, 10, 20, 30, 40, . . . 100 per cent saturation, Transfer 1-ml, portions of these solutions to test tubes of the same size as that used in the test, Add 1 ml, of freshly prepared pyrogalile-tannic acid solution to each and mix by inversion. Cover with a layer of meited paraffin and fill the remainder of the tubes with sealing wax. These standards are said to be permanent.

- 4. Neutral Methemoglobin. Dilute a little defibrinated blood (1.10) and add a few drops of freshly prepared 10 per cent solution of potassium ferricyanide. Shake this mixture and observe that the bright red color of the blood is displaced by a brownish red. Now dilute a little of this solution and examine it spectroscopically. Note the single, very dark absorption band lying to the left of D, and, if the dilution is sufficiently great, also observe the two rather faint bands lying between D and E in somewhat similar positions to those occupied by the absorption bands of oxyhemoglobin. Add a few drops of Stokes's reagent to the methemoglobin solution while it is in position before the spectroscope and note the immediate appearance of the oxyhemoglobin spectrum which is quickly followed by that of hemoglobin
- 5. Alkoline Methemoglobin. Render a neutral solution of methemoglobin, such as that used in Exp 4, above, slightly alkaline with a few drops of ammonia. Owing to the formation of alkaline methemoglobin the solution becomes redder in color and shows a spectrum different from that of the neutral solution. In this case we have a band on either side of D, the one nearer the red end of the spectrum being much the fainter. A third band, darker than either of those mentioned, lies between D and E somewhat nearer E

DETECTION OF BLOOD STAINS ON CLOTH, ETC.

 Identification of Corpuscies. If the stain under examination is on cloth, a portion should be extracted with a few drops of glycerol or physiological

³¹ Sayers and Yant Bureau of Vines Technical Paper 573 (1925) These authors describe a compact field apparatus. For gasometric determination of carbon monoxide in blood see Chapter 24

²¹ Made by dissolving 2 g. pyrogallic acid and 2 g of tannic acid in 100 ml of distilled water.

(0 9 per cent) sodium chloride solution. A drop of this solution should theo he examined under the microscope to determine if corpuscies are present

- 2 Tests on Aqueous Extract A second parties of the stain should be ex tracted with a few drops of water and the following tests made upon the aqueous extract
- (a) HEMOCHEOMOGEN Make a small amount of the extract alkaline by potas sium hydroxide or sodium hydroxide, and heat until a brownish-green color results Cool and add a few drops of ammonlum sulfide or Stokes's reageot (Appendix) and make a spectroscopic examination. Hankin has suggested a test hased upon the formation of cyanhemochromogen and the microspectroscopic demonstration of the spectrum of this compound.

(b) HEMIN TEST Make this test upon a small drop of aqueous extract ac cording to the directions given on p 483, nr, hetter, make the test upon a little material scraped from the cloth and put directly on a slide

(c) GUAIAC TEST Make this test on the aqueous extract according to the directions given on p 484 The guarac solution may also be applied directly to the stain without previous extraction in the following manner Moisten the stain with water, and after allowing it to stand several minutes, add an alco holic solution of gualac (strength about 1 60) and a little hydrogen peroxide or old turpentine The customary blue color will be observed in the presence of blood

(d) Bryzinive Reaction Make this test according to directions given on p 484

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Blood Analysis: Colorimetry and Photometry

The factors which influence the chemical composition of the blood in disease may be broadly classified as physical and metabolic. The former include those cases of retention due to alteration or destruction of permeable membranes in the excretory organs, such as the lungs, kidneys, and liver. The accumulation of nitrogenous waste products in certain forms of nephritis, and the hypercholesterolemia associated with obstruction of the biliary ducts by gallstones, are examples of retention brought about hy such processes. In the so-called metabolic diseases alteration in the chemistry of the blood may he induced by increased or diminished formation or utilization of the various constituents. Thus the accumulation of glucose in the blood as a result of metabolic derangement is an outstanding feature of diabetes mellitus. In chronic nephritus with edema (nephrosis), the high cholesterol values are considered to he of metabolic origin. Without multiplying examples, suffice it to say that changes in the blood chemistry need not be anticipated unless some condition affecting formation, utilization, or elimination is suspected. It is noteworthy, however, that the line of demarcation between these factors cannot be drawn very sharply. For example, both metabolic and mechanical factors are probably at work in renal diseases. In this connection it is important to differentiate between cause and effect. For example, the evidence that gallstones are the result and not the cause of hypercholesterolemia is not complete.

The present status of clinical chemistry is largely the result of the development of methods for the satisfactory analysis of small amounts of blood. Prior to the advent of modern micromethods for blood analysis, quantitative knowledge of metabolic processes within the body was based primarily upon analysis of the diet and urine by the classical methods of analytical chemistry. This came about because the amounts of blood required for analysis by similar methods were so large as to preclude the routine use of blood analysis for clinical purposes, except for such isolated instances as the determination of hemoglobin. Shortly after the first decade of this century, the pioneer work of such outstanding biochemists as Folin, Beaedict, Van Slyke, Myers, and their associates and pupils and many others, in developing methods for the analysis of small amounts of blood by volumetrie, colorimetric, and gasometric methods, initiated the present phase of quantitative elinical chemistry in which blood analysis has become an indispensable adjunct to the study of the functions of the body in both health and disease.

The variety of blood constituents for which quantitative methods are available and whose analytical determination is now almost routine in many hospital laboratories and elsewhere is illustrated by the accompanying table, which lists the normal concentration range for many of the major constituents of human blood, together with the chief instances in which pathological variation may be encountered. The blood of other species of animals may vary with respect to certain constituents, for example, in the rat, nonprotein nitrogen is distinctly higher, while sugar, urea, and chlorides are slightly so

For the quantitative analysis of blood and other biological material, in many cases a number of different methods are available for the determination of the same substance. This is because the aim of biochemical workers to evolve methods which are specific for the substances being determined has not always been successfully accomplished. Many methods have undergone (and are still undergoing) a process of evolution toward this goal The early blood sugar methods for example, were rela tively nonspecific and gave values of 100 to 120 mg of glucose per 100 ml of blood in normal individuals Later methods, by eliminating the effect of nonglucose reducing substances, give values lower by about 20 mg per eent or more, and presumably nearer the true glucose content. Thus the "normal" value may depend to a great extent upon the method used, and knowledge of the method employed is essential in interpreting the sig misseance of values obtained, particularly in the earlier literature The obvious value of more specific methods cannot be underestimated but for many purposes if an older and simpler but less specific method provides a guide to blood changes in pathological conditions ats usefulness continues

Because of the large number of methods which have been described in the literature, in a number of eases for the same substance, the choice of methods described in this chapter is somewhat arbitrary. The authors have selected those methods which they believe to be most generally useful and satisfactory, the selection in practically all cases being based upon personal experience or contact with hospital and research laboratories In many instances alternate methods are offered, to provide for differences of opinion concerning the relative status of methods, and for differences between laboratories in facilities available or the type of results required Blood chemical methods differ considerably in the expense or complexity of the necessary apparatus in the time required for preparation of rea gents and the performance of the analysis and in the degree of precision of the results obtained These factors must be considered in the selection of a method Certain methods for example are more adaptable than others to multiple determinations as required in large hospitals. For occasional clinical determinations where a relatively wider margin of error is admissible some of the simpler but less accurate methods may be used It is worthy of note however that the trend in modern clinical laboratories is loward more exact rather than less exact methods since only in this way can the demands of modern medicine be met

At the end of this chapter will be found a list of references to methods for the determination of blood constituents which are less commonly called for and which are not described in detail here or elsewhere in this

COMPOSITION OF HUMAN BLOOD

Constituent	Normal Runge mg per 100 ml	Pathological Conditions in Which Increases (Unless Otherwise Noted) Way Be Encountered
Total solids per cent	19 23	Anhydremua Low in hydremic plethora and
Total proteins (serum) per cent	6 5-8 2	See above Low in nephritis with edema (nephrosis)
Albumin (serum) per cent Globulin (serum) per cent	16-67	Low in nephrosis Nephrosis anaphylactic conditions malig
Fibringen (plasma) per cent	03-06	Preumonia infections Low in cirrhosis of liver chloroform or phosphorus poisoning
Hemoglobia per cent (Haden)	15 6	typhoid fever Polycythemia Low in primary and secondary anemia chlorosis
Iron as Fe Copper Total mtrogen per cent	0 03-0 25 3 0-3 7	See Hemoglobin, Vanes chiefly with proteins (albumin globulin
Nonprotein N Urea N	25-35 10-15	hemoslobin) Nephritis relampsia etc See Urea N Chronic and acute nephritis metallic poison ing cardiac failure intestinal or prostatic obstruction some infectious diseases. Rela
Une seid Creatimae Creatine Amino-acid N	2 0-3 5 1-2 3-7 5-8	tively low in nephrosis Nephritis gout arthritis celampsis Nephritis Terminal nephritis Leukemia acute yellow atrophy of the liver
Ammons N Undetermined N Gluesse	0 1-0 2 4 18 70-100	severe nephritis Terminal interstitial nephritis Eclampeia
Total fatty acids Cholesterol	290-420 150-190	Diabetes pregnancy severe nephritis Diabetes nephritis Diabetes nephritis nephrosis hiliary obstruc
Lipida phosphorus	12 14	tion pregnancy Low in permicious anomia Diabetes nephritis pregnancy In anomia low in plasma, high in cells
Total acctonn bodies (as acctone) Acctons + acctoacetic acid (as acctone) B Hydrox huty ric acid (as acctone) Bilirubin	0 8-5 0 0 3-2 0 0 5-3 0 0 1-0 25	Diabetes Diabetes Diabetes Biliary obstruction hemolytic anemias Low
CO; capacity (plasma) vol per cent	50-75†	in secondary anemia Respiratory diseases tetany Low in d abetes nephritis
CO1 content (arterial blood) vol per cent	45-50†	Respiratory diseases tetany Low in diabetes nephritis
CO: content (venous blood) vol per cent	50-60†	Respiratory diseases tetany Low in diabetes nephritis
O ₁ capacity vol per cent	16-24†	Polycythemia anhydremia Low in cardiac and respiratory diseases anemia
O1 content (arterial blood) vol per cent	15-23†	Polycythemia anhydremia Low in cardiac and respiratory diseases anenua
O: content (venous blood) vol per cent	10-15†	Polycythemia anhydremia Lowin cardiac and
Ascorbic acid Lactic acid Phenols (free)	0 8-2 4 5-20 1 2	Low in scurvy Exercise eclampsis Intestinal obstruction permicio is anemia
Chloride as NaCl milhequivalents per liter (plasma) Sulfates inorganie as S (serum) I hosphorus inorganie as I (plasma)	4:00-000 77 86 93-106 0 9 1 1 3-4	nephrats
Calcium (serum)	9 0-11 5	mg higher in children
Magnesium (serum) Sodium (serum) milhequivalents per liter Potassium (serum) millequivalents per liter lodine (104a) micrograms per 100 ml Proten lound (serum) micrograms per	1-3 300 330 130-141 16 22 4 1-5 6 8-15	Lower 12 milections of the Commission of the Com
100 ml	4-8	Hypertl yrudum and pregnancy Sometimes lux in Lypoproteinemia

[•] Figures express concentration in mg per 100 nl of wl la blood unless oil craise in heated in the first column
f rigures represent weighted averages of the observations of several investigators.

book. There is also a selected list of books and other treatises to which the reader is referred for a more comprehensive discussion than is possible in these pages of certain of the subjects considered here, particularly with reference to historical aspects and the elimical significance of laboratory data.

COLORIMETRY AND PHOTOMETRY

Introduction Many methods for the quantitative analysis of blood tissues urine, and other biological material are hased upon the produc tion of colored solutions in such a way that the intensity or depth of color so obtained may be used as a measure of the concentration of substance being determined Such use of color as an index of concentration has long been known to analytical chemists as the science of colorimetric analysis or colorimetry, and the instrument used for color evaluation is called a colorimeter These terms, while admittedly not precise, have the sanction of established usage, particularly when they refer to the very common type of color measurement in which the colored solution representing the substance in unknown concentration is brought by one means or another to exact color match with a suitable standard color Until relatively recently this was the only practical method for colorimetric analysis because the human eye is much more capable of establishing the presence or absence of exact color equivalence than it is of defining quantitative differences in color intensity in precise terms

It is clear, however, from a knowledge of the physical nature of color, that color intensity may be established in terms of the degree of light absorption at specific regions or wavelengths in the visible spectrum To take a simple example, a solution has a blue color because generally speaking it absorbs a lesser proportion of the blue components of the mixed white light passing through it than of any of the other colored com ponents Thus white light entering the solution will emerge in diminished intensity and have a preponderance of blue wavelengths. The darker the solution, ic the more intense the color the greater must be the degree of light absorption at certain wavelengths so that such light absorption may be used as a direct measure of color intensity. Analytical procedures based upon the direct measurement of color intensity in terms of light absorption at specific wavelengths are known as photometric procedures and the instrument used is called a photometer. This usage is by no means universal however, and there are many who continue to use the term "colorimetric" for methods based upon the production of colored solu tions regardless of the means by which the color intensity is established To avoid confusion it appears desirable to define a colorimetric procedure as one in which the colored solution representing the substance in unknown concentration is brought to exact color match with a standard color representing the substance in known concentration while a photometric procedure is one based upon the direct measurement of color intensity in terms of the light absorbing power of the solution at a specific region of the spectrum I blike colorimetric procedures which are limited to the visible portion of the spectrum the general principles of photometric procedures are as applicable to the absorption of radiant energy in

the ultraviolet and infrared portions of the spectrum as they are to absorption in the visible region, and increasing analytical use is being made of this fact. The use of turbidity and fluorescence for analytical purposes is discussed on pp. 533 to 537.

COLORIMETRY

As defined above, colorimetry is based upon the matching of a colored solution representing an unknown concentration of the substance undergoing analysis with a standard color representing the substance in known concentration. The substance must therefore be either colored by itself or enpable of undergoing reactions leading to the production of a color. Furthermore, the color intensity must he dependent upon the concentration, otherwise the color reaction is valueless for colorimetric purposes. A colorimetric procedure therefore involves three operations: (1) The preparation of the colored solution to represent the unknown, (2) the obtaining of a suitable standard color, and (3) color matching.

If the substance being determined is itself colored—as, for example, hemoglohin, carotene, certain inorganic 10ns, dyes, etc -the preparation of the colored solution for analysis is usually relatively simple, and may involve merely appropriate dilution or concentration of the sample to produce a color of intensity suitable for comparison against a standard. Even under these conditions, however, it is often better to separate the colored compound from possible interfering colored or noncolored material prior to estimation. If the substance must undergo a series of reactions leading ultimately to the production of a color, it is of the utmost importance to recognize that the final color intensity may be influenced to a considerable extent not only by the concentration of unknown substance hut also by the intermediate steps leading up to and including the development of the color. Such factors as the time of heating and cooling, order and rate of addition of reagents, whether the reagents are new or old, the time of standing and temperature of the solution during color development, the presence of nonchromogenic material such as neutral salts, and even the volume of solution in which the color reaction occurs, are all known to influence the final color intensity for n given amount of material in many if not all colorimetric procedures. For accurate and reproducible results, therefore, it is essential that all steps in a colorimetric procedure be carried out under conditions as earefully controlled as possible. In many cases the authors of colorimetric procedures have carefully specified the conditions for precise analysis, and these conditions should be followed without deviation.

The obtaining of a suitable standard color is obviously a most important phase of a colorimetric procedure. It may be stated without qualification that the most satisfactory standard color, and the one which should always be used for accurate results, is that obtained by treating a known concentration of the substance being determined by exactly the same procedure that is used for the unknown, at the same time, and under as nearly identical conditions as possible. Thus the final colors in standard and unknown will be due to the same substance, differing if at all only in intensity, and the many nouspecific factors already mentioned which may influence color intensity will presumably affect the standard and unknown to exactly the same extent and will not influence analytical results based upon their comparison. All colorimetric procedures must be originally based upon the use of this type of primary standard, even though secondary standards may be used later, as discussed below. This results because there is no way to predict the relationship between color intensity and concentration except in terms of the color yielded by a known concentration of the substance.

It is assumed in the use of a standard color that if the standard and unknown exactly match in color intensity they represent equal concentrations of the substance being determined. In actual practice this may or may not he true. The standard usually contains the substance being determined in relatively pure solution; in the unknown, extraneous factors may be present which modify color intensity. Substances other than the one being determined may enter into the color reaction, and results will therefore be too high; the analytical problem under these conditions is to devise either a more specific color reaction or to find methods for eliminating nonspecific interfering substances, and much of the trend in colorimetric analysis has developed along these lines Another type of interference which is less frequently recognized is the influence of nonehromogenie material present on the intensity of color produced by a given amount of chromogenie substance. This may be tested for by adding to the unknown a given amount of the pure substance, and measuring the resulting increment in color intensity If this increment is greater or less than that known to represent the added amount of substance in pure solution, and there is no possibility of loss or destruction, then factors are present in the unknown which modify color intensity per unit concentration, and suitable correction must be made. This method of using an "internal standard," as it is sometimes called, does not prove that the total color yielded by the unknown is due to the substance being determined, but it does establish whether or not this substance is capable of giving a complete color reaction under the conditions of the analysis

In some colorimetric procedures standard solutions are required containing substances which are expensive or difficult to obtain in the pure state (eg, bilirubin, and at one time, creatinine), or which deteriorate rapidly on standing or may require unavailable technical skill and apparatus for standardization (eg, hemoglobin) To provide for the routine use of colorimetric procedures based upon such standards, various "arth ficial" standards have been devised In most instances these standards consist of stable colored solutions of dyes or morganic salts, or of colored glass or gelatin The color is selected by the investigator or manufacturer to correspond as closely as possible to that representing a known amount of the substance being determined Examples of the use of artificial standards will be found in the Newcomer method for the determination of hemoglobin (p. 616), in the Benedict pierate method for the determination of urine sugar (p 923), and elsewhere In using such standards, the color in the solution being analyzed is developed by the usual procedure and then compared against the artificial standard or standards representknown amounts of material

There are many difficulties in the way of obtaining accurate results with artificial standards. Aside from the technical problem of an exact color match—and few individuals agree with one another on this point it has already been pointed out that color intensity depends not only upon concentration but also upon the technique of the analytical procedure Therefore the intensity of color corresponding to a given amount of material heing determined may vary from laboratory to laboratory Furthermore, few colorimetric procedures result in the production of a color which does not show either an increase or a decrease in intensity on standing Comparison against a simultaneously prepared standard which undergoes equivalent changes in color intensity will eliminate errors due to such changes, comparison against a stable standard will clearly give results which may depend largely upon the time of standing after color development, and careful control of this factor may therefore he necessary With few exceptions, artificial standards are satisfactory only where approximate results are sufficient, they should never be used simply to relieve the analyst of the responsibility for preparing and maintaining an exact standard solution. If artificial standards are used, they should always he checked in one's own laboratory and with one's own reagents to eliminate errors from inexact calibration, and this cheeking should be repeated at frequent intervals or when new reagents are prepared

Colorimeters. Colorimeters are instruments used to facilitate the exact matching of two colored solutions. This matching may be done in a

variety of ways, summarized as follows

1 B1 COMPARISON AGAINST A SERIES OF STANDARDS. The URLHOWN colored solution is compared by inspection with a series of color standards representing the substance being determined in known and varying con centration. The concentration of the unknown is given by the concentration of the standard which it exactly matches The colorimeter is simply a device for holding the standards and the unknown and for providing uni form conditions of illumination to facilitate exact color match. The method is simple and requires relatively inexpensive apparatus. Chief drawbacks include the labor of preparing and maintaining the standards the possibility of error due to deterioration of standards already prepared, and the fact that the range and precision of the method are limited by the number of standards available. The method therefore finds greatest application where it is known that the scope of the analytical problem is limited to results within a certain range of concentration, and highest accuracy is not required. Examples of the use of this principle are found in the colonmetric determination of hydrogen ion concentration (Chapter 1), in the Benedict pierate method for urine sugar (p. 923), and in an increasing number of "pocket," "bedside" and "field" analysis outfits available commercially for specific analytical purposes (see Fig. 121) In many instances these outfits are fitted with artificial permanent st indards, usually of colored glass or gelating to eliminate the necessity of preparing a series of standards. The use of such artificial standards has already been discussed

2 By Duplication or Colon In this method a relatively concentrated standard solution is measured into a "blank" containing the same reagents as used in the sample until the color matches that of the sample, after the volume of the stand ird his been brought up to the volume of the sample by the addition of distilled water and thoroughly mixed. The



TIC 121 SLIDE COMPARATOR FOR COLORI MITRIC COMPARISON ACAINST A SPRIFS OF STANDARDS COURTY W. A. TAYLOF CO.

Country is A Laytor Co

volume of standard solution required to prepare the duplicate is a measure of the amount of test substance in the sample. Some authors call this method colormetric thrains.



TIG 122 MYERS TEST TUBE COLORIMITER

3 By DILUTION TO COLOR MATCH Lither the unknown or the standard is diluted with water or other solvent until the two colors exactly match in intensity when compared under similar conditions of illumination and depth of solution through which the light passes. Thus if the un known must be dduted to twice its original vol ume to match a given standard, its original con centration is assumed to have been twice that of the standard, and in general if two colored solu tions differing only in concentration are brought to color match by diluting one of them, the origin nal concentration of the diluted solution is equal to the concentration of the undiluted solution multiplied by the ratio of final to initial volumes for the diluted solution Only the simplest of apparatus is required for this procedure, the color imeter may consist of two graduated test tubes (Fig 122) or cylinders of equal bore In the Sahli "Hemoglobinometer' (see Fig 161, p 612), which exemplifies this type of color comparison, the blood sample is diluted in a graduated tube to

match a permanent standard representing a known concentration of hemoglobin from the dilution required the hemoglobin content of the sample is directly determinable Dilution colorimetry 's somewhat tedious, being comparable to a volumetric titration to an irbitrary end point, but results with an error not exceeding 5 per cent may

and

be obtained with practice. A major disadvantage of the dilution procedure is that many colored solutions are affected by dilution out of direct proportion to the change in total volume owing to the influence of such factors as neutral salt concentration, acidity, or even the relative amount of solvent itself on the color intensity. Thus a blue solution of copper sulfate on heing diluted to twice its original volume will have its color intensity affected not only by the change in volume but also by the change in copper ion activity ("degree of dissociation") brought about by the dilution. It cannot be assumed that the dilution procedure is accurate, therefore, unless it has been shown to be so experimentally.

4 By Vanying the Depth of Solution through which the Light Passes This procedure is the hasis of the common laboratory visual colorimeter and is called the balancing method of color comparison. It is hased upon what is commonly known as Beer's law which for present purposes may be stated as follows 1 the intensity or density of color in a solution is determined for a particular substance solely by the number of colored particles (molecules or ions) in the light path. For example, a 1 per cent solution of a dye viewed through a solution depth of 20 mm should have the same color as a 2 per cent solution viewed through 10 mm of solution, because there are the same number of colored molecules or ions in the light path in each instance. In other words color density is directly proportional to the concentration of colored substance and the depth of solution through which the light passes. Stated mathematically, for a particular colored substance

$$D = l \times c \times l$$

where D is the color intensity or density L is the proportionality constant characteristic of the substance c is the concentration of colored substance, and l is the length or depth of solution traversed by the light heam

For two solutions of the same substance at different concentrations c_1 and c_2 , and depths of solution I_1 and I_2 .

 $D_1 = k \times c_1 \times l_1$

 $D_{\bullet} = l_{\bullet} \times c_{\bullet} \times l_{\bullet}$

If l_1 and l_2 are varied so that the two solutions have the same color intensity (i.e., $D_1 = D_2$), which is what is done in colorimeters based upon this principle, then

 $L \times c_1 \times l_1 = L \times c_2 \times l_2$

or, by transposing and canceling out L,

 $\frac{c_1}{c_2} = \frac{l_2}{l_2}$

Thus two different concentrations of the same colored substance are related inversely to the depths of solution required for color match. If one of these solutions is a standard of known concentration, C., adjusted

See pp 515 to 5.2 for a more detailed discussion of Beer s law

or

to color match with the solution of unknown concentration, C_x , by varying the depth of solution (readings R_s and R_x respectively on the colorimeter), the equation becomes

$$\begin{split} \frac{C_x}{C_s} &= \frac{R_s}{R_x} \\ C_x &= \frac{R_s}{R_x} \times C_s \end{split}$$

ie, the concentration of the unknown is guen by the readings of standard and unknown and the concentration of the standard

In using this equation, it must not be overlooked that the term concentration means amount per unit volume, hence a more exact form of the equation is as follows

or
$$\frac{\lambda}{V_X} = \frac{R_s}{R_x} \times \frac{S}{V_s}$$
$$X = \frac{R_s}{R_x} \times S \times \frac{V_X}{V_s}$$

where X and S are the actual amounts of substance present in volumes V_x and V_z , of unknown and standard colored solutions respectively, and R_x and R_z are the colormeter readings as before This form of the equation is the fundamental one upon which are based all colormetric procedures of the type described here. In most procedures the final volumes of unknown and standard are the same, and the volume factor V_x/V_z cancels out, results are then obtained primarily in terms of the actual amounts of substance present, and as a matter of fact the directions for most colormetric procedures present the taking of a definite amount rather than concentration of standard. In certain colormetric procedures (e.g., the determination of creating in blood, p. 555) the volumes of standard and unknown differ, and in such cases these volumes must be substituted in the above equation.

The value of X as obtained by the above calculation represents the amount of material in the portion of sample actually taken for color development. To express results in terms of amount per 100 ml of blood for example, X must be multiplied by 100/v, where v is the volume of blood in ml which contained the material on which the color was developed. Thus if color development in a blood sugar determination is carried out on a 2-ml portion of filtrate from the blood diluted tenfold in the preparation of the protein-free fiftrate used for the actual analysis, v in this case equals 0.2, since the 2 ml of filtrate represent 0.2 ml of original blood

In applying the above equations to colorimetric calculations, certain limitations must be noted For mechanical, optical, and analytical reasons the inverse relationship between concentration and depth of solution at color match is in general not applicable over the entire length of the colorimeter scale, which is usually about 40 to 50 mm long and graduated in millimeters. The standard is ordinarily selected to be of such

strength as to give a good intensity for color comparison at a depth of about 15 or 20 mm. It is a general rule that readings of an unknown which are less than half or more than double the reading of the standard are outside the range of application of Beer's law The amount of sample taken for analysis is selected if possible so that the expected reading will come within this range relative to the standard reading, readings outside this range are regarded as approximations only, to he used as a guide for repeating the analysis on a more satisfactor, aliquot of sample. In general, it is better to change the amount of sample analyzed so as to come within the range of the standard than to alter the standard, hecause the standard is usually so selected as to provide the most satisfactory intensity for color match, lighter or darker standards may give less accurate results If the amount of sample available is limited however, and it is known that unusually high or low values may be encountered the analysis may he saved by routinely providing several standards at different concentrations the unknown heing compared against the standard which it most closely matches on inspection

In some colorimetric procedures the range of laverse proportionality between scale reading and concentration is much less than that represented by the "half or double" rule The authors of such procedures usually specify the reliable range of readings. If the deviation from Beer's law is systematic, it is sometimes possible to establish a table of corrected values, showing the relation het een the observed readings and the amount present, thus extending the range of permissible readings. From what has been said concerning the various factors which influence color intensity, however, it is clear that such a table is highly empirical and usually reflects the conditions prevailing in one laboratory only. It should be checked at intervals for accurate results, this checking is particularly important if a table developed elsewhere is to be used

The use of light filters or of a monochromatic light source is ordinarily uanecessary in colorimetric comparisons, and mixed white light is commonly used stace if standard and unknown contain the same colored substance, at color match light transmission must be the same at all wavelengths If extraneous colored material is present (i.e., if the reagents themselves are colored), or if unknown and standard differ in hue as well as in intensity, properly selected light filters may considerably im prove the precision of readings or the range of applicability of Beer's law When artificial standards are used, uniform and reproducible illumination is particularly important because such a standard may, for exam ple, represent one concentration by daylight and a different concentration by artificial light

Colorimeters constructed to utilize the relationship between depth of solution and concentration expressed in Beer's law usually consist of (a) a source of light (mirror or built in electric lamp), (b) a pair of adjustable cups and plungers, for varying the depth of solution through which the light passes, (c) an optical arrangement for looking down through the plungers and for bringing into juxtaposition the two fields of light from the solutions being compared, to facilitate exact color match Many

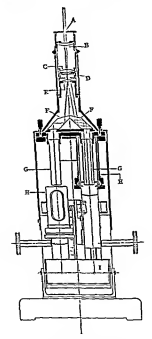


Fig 123 Diagram Showing Constitution of Duboseq Colorimeter (Bausch and Loub)

A Eye point B eye lens, C, collective D cover glass E bi prism F, rhomboid prism G, plungers, H, cups, I, mirror

different commercial models are available. The constructional details of one type are shown in Fig. 123, and other types

are illustrated in Figs 124 and 125 In the original colorimeter of this type designed by Duhosco. the cups were fixed and the plungers were raised or lowered to vary depth of solution, in modern Duhoscq type colorimeters, the plungers are fixed and the cups are adjustable. The cups usually hold about 5 ml of solution, micro-cups and plungers may be used for smaller volumes Cups with flured tops are preferred over the straightside type Some types of cups leak when in con tact with certain nonaqueous solvents such as chloroform, and are therefore unsuitable for such solvents unless sealed with a resistant cement For illumination, the most satisfactory sources are north sky light or light from an electric lamp equipped with a "daylite" filter In selecting a colorimeter, the choice should be determined largely by size and definition of the optical field and the evenness of its



ORIMETER WITH BUILT-IN LAMP IN BASE Courtesy Klett Manufac turing Co

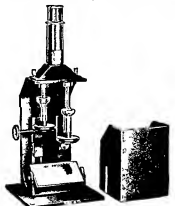


FIG 12a MICROCOLORIVETTER (BAUSCH AND I OMB)

illumination, excellence of mechanical and optical construction, and ease of operation and reading

The instruments manufactured by Klett Manufacturing Co New York Bausch and Lomb Optical Co Rochester \ \ \ and American Optical Co Buffalo \ \ \ are especi ally satisfactory in the authors experience

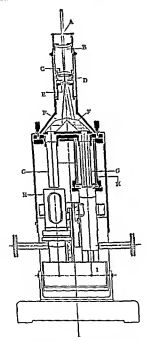


Fig. 123 Diagram Showing Construction of Duboseq Coloringter (Bausch and Lomb)

A Lye point B eye lens C collective D cover glass F bi prism F rhomboid prism G plungers II, cups, I, mirror

different commercial models are available.2 The constructional details of

one type are shown in Fig. 123, and other types are illustrated in Figs. 124 and 125. In the original colorimeter of this type, designed by Duboscq, the cups were fixed and the plungers were raised or lowered to vary depth of solution; in modern Duboscq-type colorimeters, the pluugers are fixed and the cups are adjustable. The cups usually hold about 5 ml. of solution; micro-cups and plungers may be used for smaller volumes Cups with flared tops are preferred over the straightside type, Some types of cups leak when in contact with certain nonaqueous solvents such as chloroform, and are therefore unsuitable for such solvents unless sealed with a resistant coment. For illumination, the most satisfactory sources are north sky light or light from an electric lamp equipped with a "daylite" filter. In selecting a colorimeter, the choice should be determined largely hy size and definition of the optical field and the evenness of its



Fig. 124 Klett Biocol-ORIMETER, WITH BUILT-IN LAMP IN BASE. Courtesy, Klett Manufac-

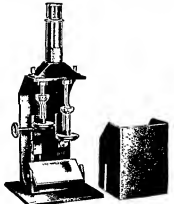


FIG 125 MICROCOLORIMETER (BAUSCH AND LOUB)

illumination, excellence of mechanical and optical construction, and case of operation and reading

The instruments manufactured by Klett Manufacturing Co., New York, Bausch and Lomb Optical Co . Rochester, N. Y., and American Optical Co , Buffalo, N. Y., are especially satisfactory in the authors' experience.

The general principles involved in the use of a Duboscq-type colorimeter are illustrated by the following experiments.

EXPERIMENTS ON COLORIMETRY

- 1. Preparation of Colorimeter for Use: (a) Librar Science. Place the colorimeter on a firm table with the mirror facing the light source. A north window or "daylite" lamp is best. Some colorimeters have a built-in electric lamp in the base; this is equally satisfactory. Look through the eyplece of the colorimeter and note the appearance of the field, focusing if necessary to bring it into sharp definition. Adjust the mirror (or the lamp in the hase) and note that the evenness and intensity of illumination can be varied at will be this adjustment. Adjust until the two halves of the field are approximately erolly illuminated, at maximum intensity. This is a preliminary adjustment only final adjustment in an analysis must always he made with the standard, at described in Exp. 2.
- (b) CHECKING THE ZEEO Place each cup, clean and dry,? on its rack heneath a plunger, and carefully rack up the cup until it comes in contact with the bottom of the plunger. Read the colorimeter scales; each should read 0.0. If they do not, locate the scale adjustment or its equivalent and adjust so that both scales read 0.0 when cup and plunger are in contact. Check by lowering the cups slightly, then raising to contact, and sgain reading. Now lower the cups and interchange them on the plungers. Again bring to contact with the plungers and read. Are the scales still in adjustment? It is common practice to mark one cup, as by wrapping a rubber hand around it, so that accidental exchange will not occur.
 - 2. Use of Colorimeter for Comparing Two Solutions. Obtain two colored solutions, a standard and an unknown, containing the same substance in slightly different concentrations. Half fill each cup with a portion of the standard, place the cups on the racks, and rack up carefully to contact with each plunger. This serves to displace air bubbles which may be trapped under the plunger. Get into the habit of routinely checking the zero at this point at the same time. Lower the cups until hoth scales read exactly 15.0. Look through the eyeplece. Adjust the light source carefully until the two halves of the field are as evenly illuminated as possible. Theoretically, the reading on each side is now 15 0; actually, there is as much error in this adjustment as In the subsequent reading of an unknown; therefore for precise results the solution on one side should now be read against the other side ("matching the standard against itself"). Leaving the left-hand cup set, lower the righthand cup slightly to throw the two halves of the field out of balance, then raise the cup slowly while looking through the eyeplece, until the field appears exactly even. Note the right-hand scale reading. Repeat this process three or four more times, and average the readings. This is the actual reading of the standard that is used in the calculations.

When the reading of the standard in the right-hand cup has been established, remove this cup, discard its contents, and place a portion of the unknown in the cup, iloiding the cup in the hand, run it up and down hreffly on the right-hand plunger, thus rinsing cup and plunger with the unknown. Discard the cup contents and repeat the runsing with a fresh portion. Finally half fill the cup with fresh unknown and replace it on the cup rack. Run it up carefully to contact with the plunger, to displace air bubbles, then Jower it

¹ Use a fresh piece of soft lintless paper dens paper or Kleenex) to wipe the hottom of each cup drj. Do not use a towel or handkerchief for this purpose

until inspection through the eyepiece indicates color match. Make the reading, and repeat three or four times as described for the standard, averaging the readings. The average result is the reading of the unknown.

From the readings of standard and unknown, and the known concentration of the standard, the concentration of material in the unknown is calculated as follows:

$$\frac{\text{Reading of Standard}}{\text{Reading of Unknown}} \times \frac{\text{Concentration}}{\text{of Standard}} = \frac{\text{Concentration}}{\text{of Unknown}}$$

Note that in this calculation the readings of standard and unknown are hoth made on the same scale, i.e., the right-hand scale. The standard in the left-hand cup is set at 150 (or 200, or whatever value is specified) merely to establish a satisfactory color intensity for comparison purposes, and its reading does not enter into the calculations. As a matter of fact, some types of colorimeter do not have more than one scale. It is true that various other methods of reading a colorimeter have heen proposed, but these are all subject to more error than the procedure described here

In making colorimeter readings, it is good practice to rest the eyes frequently by looking up from the colorimeter, because the eyes tire easily and color discrimination becomes less acute. Make four or five independent readings and average them to minimize error, if one reading in the series is obviously quite different from the others it may be discarded in averaging. Best results are obtained if alternate readings are approached from above and below, and the hand removed from the instrument between readings, thus minimizing the tendency to return mechanically to the same reading.

- 3 Influence of "Personal Error" on Colorumetry Repeat the above experiment, but after you have read the standard against itself, have another individual make independent readings, and compare results How closely do you agree? Now read the unknown against the standard, and calculate the concentration of the unknown from your readings. Have the second person likewise read the unknown and calculate results from his readings. Is any difference in opinion concerning the readings reflected in the final results? In general, it is necessary for each analyst to make his own readings of both standard and unknown, in this way, differences of opinion as to precise color match do not influence results.
- 4 Beer's Lan Obtain a series of aqueous methylene blue solutions containing 0 004, 0 005, 0 100, 0 200, and 0 030 g. per liter respectively of the dye Select the 0 010 g. per liter solution as a standard, and set at 15 mm. Match the standard against itself, making the readings as described in Exp. 2, and then read each of the other solutions against this standard. On a sheet of cross section paper, plot the colorimeter readings on the 3 axis against the concentration on the x axis. Draw a smooth curve between the points. Now calculate by the use of Beer's law what the various readings should be for the determined reading of the standard. Plot these theoretical readings on the same graph, and connect the points hy a smooth curve. Do the two curves overlap completely or do they diverge at any points? To what extent does this dye follow Beer's law under these conditions? How could you obtain accurate results in a colorimetric procedure which is known not to obey. Beer's law?

Another way of plotting results is to plot the theoretical reading for a given concentration against the actual reading. A straight line indicates adherence to Beer's law. It is the general practice to obtain results by calculation based on Beer's law only if the unknown reading is not more than double or less than half the standard reading. Unknowns falling outside this range are repeated with more or less of the sample for analyses.

5 Influence of Analysical Procedure on Color Intensity Into each of two small flasks place 20 ml of standard ammonium sulfate solution contain ing 0.2 mg of nitrogen (seep 549) To one flask add 2 ml of Nessler's reagent 'drop by drop, from a buret (Because of its poisonous nature, 'never use a plipet with Nessler's reagent) Set aside, and to the second flask add 2 ml of the Nessler reagent, as rapidly as possible. Mix hy lateral shaking and set aside After 10 minutes standing, read one of the two solutions against Itself in the colorimeter, and then read the second solution against the first. Do the readings indicate that the two solutions have the same concentration of ammonia? What must be done to ensure uniform results?

PHOTOMETRY

Photometry, in the sense in which the term is understood in analytical chemistry, consists in the measurement of the light-transmitting power of a solution in order to determine the concentration of light-absorbing material present For purposes of simplicity the term light is used in place of the more inclusive term radiant energy. It must be understood that the principles upon which photometry is hased are as applicable to the transmission of energy in the ultraviolet or infrared portions of the spectrum as they are to transmission in the more commonly employed visible, or colored, spectrum (see Fig. 287, p. 1249 for a chart showing the spectral distribution of radiant energy)

The ability of a solution to transmit light is known as the transmittance, T, of the solution Strictly speaking transmittance is defined as the ratio of the intensity, I_2 of the light emerging from the cell containing the solution to the entering or modent light intensity, I_1 , or

$$T=\frac{I_2}{I}$$

For photometric purposes however it is neither practical nor necessary to measure the incident light intensity and to define transmittance in this way. In a photometric procedure there is always a certain proportion of nonspecific light loss during passage through the solution due to reflection from glass surfaces scattering and absorption by the solvent, reagents and even possibly by contaminating traces of the substance which is being determined. Any of these will obviously influence a single measurement but should be constant from one measurement to another, and may therefore be balanced out by defining transmittance in relative terms. Thus if we let T_{sol} be the (over all) transmittance of a solution containing a light-absorbing material (the solution and the T_{sol} be the (over all).

⁴ See Annendi

^{*} See Chapter 31 for a discussion of the chemistry of hessler a reagent and the reaction with ammonia.

transmittance of a reference solution (the solvent), usually the colorless solvent or reagent blank hoth solutions heng examined under equivalent conditions of wavelength, incident light intensity, and depth of solution, then the ratio of the transmittance of the solution to that of the solvent (or reagent blank) is equal to T_s, the transmittancy of the sample, 1 e,

$$T_* = \frac{T_{soln}}{T_{roln}} = \text{Transmittancy}$$

In this way neither the intensity of the incident light nor the nonspecific light loss need he determined, and furthermore contaminating traces of the substance heing determined, in reagents which are themselves colored, do not interfere in an analysis. The change in transmittance due to the presence of the substance is determined solely by the increase in light absorption above a level which is arbitrarily taken as zero. It is this possibility of ruling out error from blank light absorption which represents an outstanding advantage of photometric analysis over colorimetric analysis with the Duboseq type colorimeter, where this cannot be carried out without the use of correction factors.

Transmittancy, T_* , is thus a relative measurement and is always less than 100 if light-absorbing material is present It may be expressed numerically either as a decimal fraction or in terms of per cent, e.g., a transmittancy of 0.65 or 65 per cent. A more satisfactory way of expressing the transmittancy of a solution is in terms of its negative logarithm, $-\log T_*$, known as the absorbancy, A_* also designated as the optical density, D_* , or the extinction, E_* , of the solution

$$A_{\bullet} = -\log T_{\bullet} = \log \frac{1}{T_{\bullet}} = \text{Absorbancy}$$

The utility of this hasis for defining light absorption, particularly for photometric purposes, will be presented subsequently

Determination of Transmittancy. The light transmittancy of a solution is determined by the use of an instrument known as a photometer Many varieties and designs of photometers have been described and are available commercially, regardless of design, the principle upon which all analytical photometers operate is fundamentally the same, and may be described as follows Light of suitable wavelength is allowed to pass through a reference solution, usually the colorless solvent or reagent blank, held in a container of fived dimensions known as a cuvette. The intensity of light emerging from the reference solution is established at an

iii It is felt by some that a distinction between transmittance and transmittancy as defined here is relatively unimportant Until standardized nonmerclature in this field becomes more universally adopted one max find some authors or publications using one term and some the other when referring to no totometrue measurement.

Some investigation particularly, in Furopean laboratories use the natural logarithm rather than the common logarithm is — In Trather than — log T Tle choice is immaterial for photometric purposes but may lead to confusion in applying data obtained elsewider Tle relation between these two ment of so for pression is as follows.

arbitrary value by any of the various methods described below, this value usually corresponding to a reading on the photometer scale of 0 optical density or 100 per cent transmittancy. The reference solution is then replaced by the solution whose transmittancy is to be determined, held in the same or a similar cuvette, and the emergent light intensity measured relative to that established for the reference solution, this relation gives the transmittancy of the solution under examination

The intensity of light emerging from a solution may he established by either visual, photographic, or photoelectric (or equivalent) means, of these the last is most common, most accurate, and has largely displaced the others In a visual photometer, the emergent light beam is compared in intensity with a parallel reference beam of similar properties which is of arhitrary and adjustable intensity. The adjustment required to bring the reference beam to the same intensity as that emerging from the solution under examination is the measure of the emergent light intensity Measurements are therefore influenced by the acuity of visual color intensity discrimination as in visual colorimetry In the photographic plate method the intensity of action of the emergent light on a photographic plate is compared with the action of light of relatively known and adjust able intensity under similar conditions. This procedure is tedious and has been used in the past largely for measurements in the ultraviolet and infrared portions of the spectrum, where the eye is insensitive Much of the earlier data in the literature concerning light absorption in these regions of the spectrum is based on this method, but it has been super seded almost entirely by the use of light-sensitive devices such as the photoelectric cell or its comvalent

Light intensity is determined photoelectrically by using photoelectric cells or similar light-sensitive devices which produce an electric current in proportion to the intensity of light striking their active surfaces Two types in common use are (1) the photovoltaic cell ("plate type," "recti fier" or "barrier layer" cell) and (2) the photoemissive tube A photovoltage cell consists essentially of a metal plate coated with light-sensitive material (selenium, cuprous oxide) which is in turn coated with a thin transparent film of a metal such as gold or copper Light passing through the transparent film sets up a flow of electrons in one direction ("recti, fied") which establishes a potential difference between the two poles of the cell, and causes a current to flow if the cell is in a suitable electric circuit Because of the electrical characteristies of photovoltaic cells the current is not suitable for amplification but for ordinary light intensities it is sufficiently large to be registered on a microammeter or low sensitiv ity galvanometer, and many types of photometers employ such cells Photoemissive tubes or phototubes are either evacuated or gas-filled tubes similar in appearance to radio tubes and containing a plate coated with some substance which emits electrons when light strikes it the electrons traveling to a suitable anode under proper conditions. The in tensity of incident light thus determines the flow of electrons through the tube, and hence the current in an external circuit. This current is very small but may be readily amplified and photometers employing such tubes usually have an amplifier circuit. An advantage of phototubes 15

that they may be obtained with wide ultraviolet and infrared sensitivity, which is not the case with the photovoltaic cell.

With photosensitive devices, change in current output is used as a measure of change in light intensity. If the current output is adjusted to an arbitrary value with the reference or blank solution in place in the photometer, then the transmittancy of an unknown solution is given by the ratio of current output for this solution as compared to that for the reference solution. It is assumed that the current output is strictly proportional to the light intensity; with well-designed photometers thus is usually sufficiently true over the range of light intensity for which they are used. If it is not the case, there may be apparent deviation from the theoretical relation between transmittance and concentration as defined by Beer's law (see p. 516), and such photometers are usually worthless for analytical purposes Since the transmittancy is measured in a photoelectric photometer in terms of current output rather than of actual light intensity, it has been proposed that in such instances the term photometric density be used for the value of - log T, rather than optical density. From a practical point of view this distinction is unimportant and will not be used here, but it serves to emphasize the point that the numerical value of the optical density for a given substance may be considerably influenced by the characteristics of the photometer used for the measurement.

Light absorption at specific wavelengths is an intrinsic property of many substances, eg. hemoglobiu and other colored compounds in the visible region of the spectrum, and many other compounds in the ultraviolet and infrared portions of the spectrum. It may therefore be used to characterize such substances, just as other physical constants are so used, in addition to its use in photometric analysis. The extinction at a specified wavelength for a unit amount of a particular substance in solution is known as the specific extinction or extinction coefficient; the amount is frequently defined as that present in a 1-cm layer of solution containing 1 per cent of the substance (symbolized by E10). If the concentration is expressed on a molar basis, the term molar (or molecular) extinction is used These constants are of more value for characterizing the substance in terms of its optical properties than for actual use in photometrie analysis, but if they are sufficiently reproducible they may be used for the latter purpose, since they define the relationship between concentration and light absorption for the particular substance under specified conditions.

Beer's Law. The transmittance of a solution containing light-absorbmaterial depends upon (a) the nature of the substance, (b) the wavelength of the light, and (c) the amount of light-absorbing material in the light path, this latter depending in turn upon the concentration of substance and the depth of solution through which the light pa-ses. The relation between these various factors was first clearly established for colored solutions by Beer, and hence is known as Beer's law, in-o as the Bougner-Beer or Lambert-Beer law. This relation may be expressed as

⁷ For a fuller discussion of the laws of Bouguer and Beer, see Mellon: Analytical Absorption Spectroscopy, New York, John Wiley & Sons, Inc., 1950.

or

follows at a given wavelength,

where T_i is the internal transmittance, L is a constant characteristic of the substance, l is the length or depth of solution through which the light passes, and c is the concentration of light-absorbing material T_i is defined as the ratio l/I_o , where I_o is the radiant energy entering the sample and I that incident upon the second surface of the cell (see Fig 126) The equation is exponential because of the particular characteristics of light

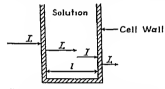


Fig 126 Schematic Diagram Showing Derivation of Transmittance (Ratio I_r/I_1)

Internal transmittance T_t is the ratio I/I_s l is the length or depth of the absorbing path in the sample

absorption (see texts on physics for details), it may be converted to the more common logarithmic form by taking the logarithm of both sides as follows

$$\log T_i = -l \times l \times c$$

 $-\log T_i = l \times l \times c$

These equations relating transmittance concentration, and depth of solution at a given wavelength, are the fundamental ones upon which photometric analysis is based.

As mentioned on p 512, for photometric purposes it is more practical to define transmittance in relative terms ie the ratio of the (over all) transmittance of the solution T_{-i+} to that of the solvent (or reagent blank), T_{-i+} , equal to T_- the transmittancy T_- is not precisely equal to $(T_-)_{-i+}/(T_-)_{-i+}/(T_-)_{-i+}$, but with end plates having a refractive index not greater than 15, the error is usually negligible

Relation between Transmittance and Concentration. If the transmittances (in practice, transmittance, T, values) of a series of solutions of a particular substance in various known concentrations are determined, at a particular wavelength and constant depth of solution (the issual conditions in a photometric analysis), the resultant data relating transmittance to concentration may be plotted in any one of four different ways as shown in Fig. 127. If the transmittancy is plotted directly against concentration on ordinary cross-section paper, the cure

of Fig. 127A is obtained To obtain a straight line rather than a curve, and thus permit the accurate establishment of the relationship hetween transmittancy and concentration at only one or two concentrations instead of the many required for a curve, advantage is taken of the fact that in accordance with Beer's law the equation for the curve is:

$$\log T_* = -k' \times c$$

the constant depth of solution permitting the combination of l and k into k. Thus if the logarithm of the transmittancy is plotted against concentration (Fig. 127B), or if transmittancy is plotted on the logarithmic

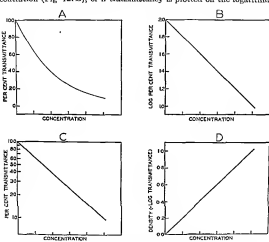


FIG. 127. VARIOUS WAYS OF SHOWING GRAPHICALLY THE RELATION BETWEEN LIGHT TRANSMITTANCE AND CONCENTRATION IN A PHOTOMETRIC PROCEDURE.

A, Per cent transmittance plotted directly against concentration, B, Log per cent transmittance against concentration. C, Per cent transmittance against concentration on semilograthmic paper. D, Density (- log transmittance, or 2 - log per cent transmittance) against concentration.

axis of semilogarithmic paper (Fig. 127C), straight lines with negative slopes will result. Finally, if the Beer's law equation is written as follows:

$$-\log T_* = L' \times c$$

and the values of $-\log T$, plotted against c, a straight line with a positive slope results (Fig. 127D).

Of these four methods for relating transmittancy to concentration, the last is to be preferred. The quantity—log 7, has niready been defined as the optical density, D, of the solution. Where Beer's law is applicable, the optical density is directly proportional to concentration, or

$$D = 1' \times c$$

This represents the simplest possible relation between light absorption and concentration, and it is the major advantage obtained by expressing transmittains, in terms of optical density rather than as per cent

The established relationship between transmittanes and concentration for a particular analytical procedure is known as the ealibration for that procedure. If this relationship is shown in the form of a graph, as in Fig. 127, the graph is called the calibration curve. Every analytical procedure requires a separate calibration, over a specified range of concentration and at a given wavelength and depth of solution, and in general the calibration established for one photometer is not applicable to another photometer, even of the same make The most accurate way to establish the ealthration for a particular procedure, and the one which should be used whenever possible, is in terms of the reading of a simultaneou.ly prepared standard (or, in some instances, series of standards) for that procedure, just as in accurate visual colorimetry, and for the same reasons In practically all of the common colorimetric or photometric procedures the substance being determined is subjected to a series of reactions leading to the final production of a color which is used as the basis for estimation The many factors aside from concentration which influence the final color intensity in most if not all colorimetric procedures have been discussed on p 501 in connection with colorimetric analysis they are of equal if not greater importance in photometry. Furthermore since photometric measure urements are ordinarily made individually and in a sense "against the instrument itself," rather than by comparison against a standard color variations in mechanical optical, and electrical properties of the photom eter may likewise influence the calibration. Thus the calibration for \$ particular procedure may vary from day to day, from one photometer to another, and from laboratory to laboratory If calibration is established in terms of the reading of a simultaneously prepared standard at the time the analysis is carried out, the various factors other than concentration which may influence the actual readings exert the same effect on the standard and on the solution being analyzed, and thus do not influence the retative evaluation of one in terms of the other

To calibrate a procedure in terms of the reading of a simultaneously prepared standard advantage is taken of the fact that optical density (or extinction) is directly proportional to concentration where Beer's law is applicable, as it is to practically all photometric procedures under the proper conditions. Therefore for two solutions of the same substance at different concentrations at the same wavelength and depth of solution the relation between their respective densities, D_1 and D_2 , and concentrations, c. and C_2 , is as follows

 $\frac{D_1}{D_2} = \frac{c_1}{c_2}$

If one of these two solutions is a standard of known concentration and the other is of unknown concentration, then if their respective densities are measured, the concentration of the unknown is given by the calculation

$$Concentration \ of \ Unknown = \frac{Density \ of \ Unknown}{Density \ of \ Standard} \times \frac{Concentration \ of \ Standard}{Standard}$$

Thus the concentration of an unknown in a photometric procedure which shows agreement with Beer's law may be calculated from the determined density and the density of a known standard, and this method of obtaining results in a photometric acalysis is used where applicable for all of the photometric procedures described in this chapter and elsewhere in this book.

The calculation is obviously quite similar to that used for visual colorimetric analysis with the Duboseq type colorimeter (see p. 505), except for the direct instead of inverse proportionality hetween concentrations and readings. It is subject to the same limitations concerning the use of concentration rather than amount, if amount is used in the above equation rather than concentration, the unknown and standard mushave the same total volume, otherwise the right-hand side of the equation is multiplied by Y_x/V_s , where V_x is the total volume of unknown and V_s the volume of standard. To express results in terms of 100 ml of blood, per liter of urine, or on some other arbitrary basis, the obtained result must be multiplied by a factor representing the relationship between the actual amount of sample used and the desired basis, as for colorimetric calculations

To use this method of calculating results with photometers whose scale reads only in terms of per cent transmittancy, it is occessary to change the transmittancy value into its equivalent optical density. This may be done using a table of logarithms, or more simply by reference to the accompanying table, which gives the value of the optical density for all values of per cent transmittancy. Thus in an analysis, if the per cent transmittancy of standard and unknown are determined, readings are converted ioto density values by reference to the table, and results calculated as described above. If the photometer scale gives the density values directly, or reads in units which are proportional to density, the scale reading is used directly in the calculations. This is obviously more convenient, and a photometer scale of this type is to be preferred over one which is in terms of per cent transmittancy only.

Calculation of photometric results by the method described can of course be used only over the range of concentration where Beer's law is valid, i.e., where there is a linear relationship between optical density and concentration. For practically all photometric procedures in common use conditions of wavelength, depth of solution, and concentration may be so selected that Beer's law is obeyed over the range of concentration apt to be encountered in an analysis. It is important in describing the details of a procedure to define these conditions, as has been done for the photometric procedures described in this chapter and elsewhere in this hook.

Relation between Transmittance (T_{\bullet}) and Optical Density (D)

T. (%)	D	T. (%)	D	T. (%)	D	T. (%)	D
100	0 000	75	0 125	50	0 301	25	0 602
99	0 004	74	0 131	49	0 310	21	0 620
98	0 000	73	0 137	48	0 319	23	0 638
97	0 013	72	0 143	47	0 328	22	0 658
96	0 018	71	0 119	46	0 337	21	0 678
90	0 022	70	0 155	45	0 347	20	0 699
91	0 02"	69	0 161	-41	0 357	19	0 72
93	0 032	68	0 168	43	0 367	18	0 74
92	0 036	67	0 174	42	0 377	17	0 77
91	0 011	66	0 181	41	0 387	16	0.79
90	0.046	65	0 187	40	0 398	15	0 82
89	1c0 0	61	0 194	39	0 409	14	0 85
88	0 056	63	0 201	38	0 420	13	0 884
87	0 061	62	0 208	37	0 432	12	0 92
86	0 066	61	0 215	36	0 444	11	0 95
85	0 071	60	0 222	35	0 456	10	1 00
84	0 076	59	0 229	34	0 109	0	1 01
83	0 081	58	0 237	33	0 482	8	1 09
82	0.086	57	0 214	32	0 495	7	1 15
81	0 092	56	0 252	31	0 509	6	1 22
80	0 097	55	0 260	30	0 523	5	1 30
79	0 102	54	0 268	29	0 538	4	1 39
78	0 108	53	0 276	28	0 552	3	1 52
77	0 114	52	0 284	27	0 569	2	1 69
76	0 119	51	0 292	26	0 585	1	200

On photometers equipped with a linear scale reading from 0 to 100 D corresponds to the value of $2 - \log G$ where G is the galvanometer or microammeter realing relative to an initial set ting at the 100 mark

For a few procedures of which the determination of blood creatinine by the alkaline pierate reaction (p 555) is an example it is found that Beer s law is not followed under any analytical conditions In such cases a calibration curve must be constructed and results obtained by reference to the curve as described below for the most accurate results such a curve should be constructed with each series of analyses unless it is found to be highly reproducible Instances of deviation from Beer's law are fortu nately very rare It is more common to find an apparent deviation from Beer's law when agreement is expected. This is usually the fault of the photometer In most instances agreement with Beer's law can be expected only when essentially monochromatic light is used particularly when photovoltaic cells with their inherent varying spectral sensitivity are used in the photometer If a wide rather than a narrow portion of the spectrum is used for measurement as is unfortunately the case with some types of photometers or if the photocell circuit is such that current output is not proportional to light intensity the relationship between optical density (or $-\log T$.) and concentration will not be linear, and results must he based upon a calibration curve Photometers with such characteristics are therefore undesirable as a basis for accurate photometric analysis, hecause they do not permit the analytical precision associated with the application of Beer's law to the procedure.

Another commonly used method for obtaining results in a photometric analysis is based upon the use of previously prepared graphs of the type shown in Fig. 127, or of tables based upon the graphical data. The graph or table is established for a particular procedure by determining the transmittancy values for a sufficient number of solutions of known and varying concentration with respect to the substance being determined. In future analyses, the traosmittancy of the solution of substance in unknown concentration is determined, and its concentration is then found from the graph or table, without running a standard at the same time ("colorimetry without standard solutions") It is not occessary that the color reaction show agreement with Beer's law, since the relation between photometer reading and concentration is established empirically; this type of procedure therefore finds its greatest use where for one reason or another there is lack of agreement with Beer's law. An analogous procedure in instances where Beer's law is valid is to establish the density of a known standard and to use this value in future analyses, i.e., the concentration of the unknown is obtained by multiplying its determined density by a factor representing the established relationship between the standard and its density (see the photometric determination of hemoglobin, p. 610, for examples). The extinction coefficient of a substance may also be used in a similar nav.

It is assumed in the use of such previously obtained calibration data that a particular transmittancy or density will always represent a particular coocentration in an analysis In practice this may or may not he true. The many factors which influence color intensity aside from concentration have already been emphasized. Even for substances such as hemoglobin in which the light-absorbing power is an integral property of the molecule itself, variations in environmental conditions or in the photometer itself may influence readings. Thus the use of a previously prepared calibration curve may be at best only an approximation, and gross errors are known to have resulted from its use. Colorimetry without standard solutions does not exist; the use of a previously prepared calibration curve simply represents a decision that the standard is to be prepared and read at one time and the unknown at some other time and possibly under different conditions, rather than that standard and unknown are to be prepared and read under the same conditions. It may be stated without equivocation that the greater desirability of the latter procedure has never been seriously challenged.

It is true that in some instances, as where the standard substance is difficult to obtain or maintain stable in solution, or the scope of the analytical problem permits the sacrifice of accuracy to convenience, a calibration curve may satisfy analytical requirements If such a curve is used, it should be constructed in one's own laboratory using the reagents and photometer which will actually be employed in the analysis Calibration data obtained from the literature or from the manufacturer of the pho-

tometer should never be used without checking and this checking must be repeated at frequent intervals if satisfactory results are to be expected particularly if there has been a change in the reagents or photometer The analyses must be carned out with rigorous control of the various step involved reproducing as far as possible the conditions under which if curve was constructed. Only in this way can the occurrence of sense errors be prevented.

Relation between Transmittancy and Wavelength The relative between the transmittancy (and hence the optical density) of a solution containing light-absorbing material and the wavelength of light passification of the solution is given by the so-called absorption spectrum of the substance. The absorption spectrum is established quantitatively the measuring the transmittancy for a particular concentration and depth solution at various wavelengths, and plotting the results in the form of

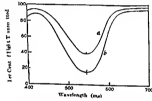


Fig 128 A Relatively Simple Absorption Spectrum

Curve b represents a concentration twice as great as for curve a. The dotted incin licates the wavelength of maximum gensitivity for photometric measurement.

enric relating transmittancy or optical density (the latter is preferred to wave-length. An example of a relatively simple absorption spectrum is shown in Fig. 128, more complex curves are frequently found. The absorption spectrum of a substance is usually characteristic of the substance and may serve for identification as well as furmishing information of analytical value. The application of absorption spectra is not limited to the visible region of the spectrum but may be applied equally well to characterization of the ultraviolet or infrared absorption of many substances.

For a particular substance absorption curves at different concentrations will be generally similar in shape but will differ in their positional properties of the properties o

the cause of color, since when white light cantaining all wavelengths passes through the solution the emergent light cantains a greater proportion of some wavelengths than of others, and hence appears colored

As the concentration of substance in solution is increased (Curve b relative to Curve a) there is usually increased light absorption at all wavelengths where any light is absorbed at all but this increase is usually greater per unit change in concentration at some wavelengths than at others. Thus the wavelength used may determine the relationship hetween concentration and optical density at a canstant depth of solution (i.e., the numerical value of I in the Beer's law equation), and this is frequently a major factor in the choice of the praper wavelength for photometric measurement.

Maximum sensitivity in a photometric procedure is obtained at the wavelength where there is the greatest change in optical density or trans mittaney per umit change in concentration. Most often this is the wavelength of maximum light absorption or minimum in the transmittiney curve (dotted line. Fig. 128). Occasionally, bowever, as for example in a procedure where the transmittancy curves for reagent and colored compound overlap considerably, the highest sensitivity may be obtained at a wavelength on one side of the absorption maximum. Thus, a knowledge of the absorption spectrum of both reagent and colored compound (the latter at two or more concentrations) is necessary for selection of the optimum wavelength.

Maximum sensitivity is not per se the chief consideration in the selection of the proper wavelength for photometric measurement. From an analytical point of view the most satisfactory wavelength is the one which at a given depth of solution shows agreement with Beer's law over as wide a range as possible of the concentrations apt to be encountered in an analysis and which permits this range to be read within the most accurate region of the photometer scale. The most accurate region of the scale corresponds to densities between about 02 and 10 (60 to 10 per cent transmittancy) Readings outside this range represent solu tions which are either too light or too dark for the most accurate measurement thus at 95 per cent transmittancy an absolute error of 0.5 per cent in the transmittaney measurement corresponds to a 10 per cent error in an analysis, and at the other end of the scale for dark solutions unit change in transmittancy represents a disproportionately large change in concentration. The sample should therefore read between the scalo limits specified if maximal accuracy is to be abtained.

To fulfill these requirements in the case of the Fohn Wu hlood sugar method, for example (see p. 508) a wavelength may be selected which represents a very low sensitivity, so that the normal glucose standard representing 100 mg per cent blood sugar will have a low light absorption. This would permit rading blood sugar values well above normal which is the usual direction of change in this procedure under the same conditions. If the usual direction of change is below the normal value as in himoglobin determinations conditions are selected such that the normal sample has a high light absorption. In general, the adapt to photometric measurement a procedure which was originally developed for visual

colormetry and hence may yield a more intense color than is required for photometric measurement, and in a larger volume of solution, it is better to modify the light absorption. This may be done by selection of a suitable wavelength or use of a small depth of solution rather than by such procedures as taking a smaller sample or diluting the final color, since these may scriously affect the accuracy. However, the greater sensitivity of photometric measurement, and the possibility of using very small volumes of solution, is obviously conducive to the development of microanalytical methods, of which many have been and are heng developed, and this represents an important contribution of photometry to analytical chemistry.

Other factors which may influence the choice of wavelength for photometric measurement include the possibility that agreement with Beer's law will be found to be more satisfactory over a wider range of concentra tion at one wavelength than at another, or that the color will he more stable when exposed to light of one wavelength than to that of another Even if such selection entails a decrease in sensitivity, the enhancement of the analytical value of the procedure may make the wavelength of lesser sensitivity the one of choice. If two or more light-absorbing substances are present together, and it is desired to measure the change in optical density related to variation in amount of only one of these substances, it is semetimes possible to select a wavelength at which there is only minimal light-absorption by extraneous material, and thus prevent such material from interfering significantly in an analysis Such photometric separation is rarely complete, it is usually better to effect preliminary analytical separation, or to include the light absorption from extraneous material in the blank solution used for the initial setting of the photometer

Knowledge of the complete spectral characteristics of a light-absorbing compound is of fundamental importance in defining the conditions under which satisfactory photometric analysis is possible. Wherever possible such information should be made available in the description of a photometric procedure, as has been done in many instances for the various photometric procedures described in this chapter and elsewhere in this book. The detailed description of the spectrophotometric characteristics of the Nessler reaction with ammonia, given in Chapter 31, illustrates the applicability of absorption spectrum data to photometric procedures.

Light Filters and Filter Photometers. The wavelength at which photometric measurements are made may be established either by the use of light filters or by the production of a complete spectrum and isolation of the desired portion. Instruments based upon the first principle are known as filter photometers, those based upon the second principle are called spectrophotometers (see p. 531). Most types of photometers in current use are filter photometers, though spectrophotometers are now extensively used even for routine analyses. Filter photometers, however, are less expensive than spectrophotometers require less technical skill in operation and if well designed are just as satisfactory for most analytical purposes particularly in the visible region of the spectrum. Except for special instances spectrophotometers must ordinarily be used for photometry in the ultraviolet and infrared portions of the spectrum.

Light filters commonly consist of selected glass (or sometimes dyed gelatin) which is capable of transmitting light over a limited portion of the spectrum only. Thus by placing such a filter in the light path of the photometer, measurements may be made in the spectral region corresponding to the transmittance range of the filter. Various filters differ principally with regard to (a) the spectral region of light transmittance, and (b) the width of the transmitted band. By suitable selection of various types or combinations of glass, it is usually possible to obtain a filter whose transmittance is limited to almost any desired portion of the spectrum. A selection of such filters is illustrated in Fig. 129. Suitable filters may be obtained from the manufacturers of the photometer, or they may be constructed in the laboratory if suitable glass is available It is customary to designate a filter in terms of the wavelength of peak

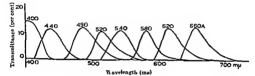


Fig. 129, LIGHT TRANSMITTANCE OF SELECTED LIGHT FILTERS.
Courtest, Rudicoa Co

transmittance, thus a filter called "No. 540" or "No. 54" has its peak transmittance at a wavelength of 540 m μ This practice is not universal, nor is the wavelength designation always accurate, and when a filter of any kind is used in a photometric procedure the wavelength of peak transmittance should be stated also If this information is not known, it may usually be obtained from the manufacturer of the filter or glasses used. When the wavelength required for a particular photometric procedure is stated in the description of the procedure, as for example, "at 540 m μ ," this corresponds to the use in a filter photometer of a filter with a peak transmittance at this wavelength

The most satisfactory filters are those which transmit as narrow a spectral region as possible, since this represents an approach to truly monochromatic light. A good filter for photometric purposes will show a transmittance of about 85 per cent or more of the total light transmitted over a spectral width of 30 to 60 mg or so, centered around the wavelength of peak transmittance. This information concerning a filter may be obtained from its transmittance curve, as illustrated in Fig. 120. Filters with a broader range of transmittance are in general unsatisfactory, because they may result in apparent deviations from Beer's law, as discussed on p. 520.

In recent years interference filters (metallic films on glass or fused quartz) have come into use, such as, for example, those made by Baird, Farrand, and others. The Farrand filters consist of exaporated thin layers of dielectric material between semitransparent metallic films on glass.

Interference filters are sometimes called monochromatic filters because of their band pass of only 20 m μ or less much narrower than the usual glass or gelatin filters. These filters, however, are not monochromatic in the sense that hight of a single wavelength is monochromatic, since they transmit a narrow range of wavelengths rather than a single wavelength



Fig 130 Hellige Chromatron Photoelectric Colorimeter Courtesy Hellige Inc

Interference filters have a high transmission being about 40 per cent—more than twice that posble with ordinary filters of equal band pass

Representative types of filter photometers commercially available are illustrated in Figs 130, 131, and 133 Many other types are on the market,* some of which are doubtless as satisfactory as those illustrated The Hellige "Chromatroa (Fig 130) is an example of the single photocell type of photometr with huilt in meter, the scale of which reads linearly from 0 to 100 and hence gives transmittancy in per cent. This instrument is designed for use with rectangular pyres.

cuvettes The light source is mounted in a self focusing housing and is oper ated below its rated wattage either from two dry cell batteries or from an A C line through a huilt-in constant voltage transformer The filters are mounted in a rotating disk close enough to the photocell to avoid interfer ence from stray light The photocell is hermetically sealed in an mert at mosphere In operation, the instrument is adjusted to a scale reading of 100 with water and a suitable filter in place the water is then replaced by the solution under examination and the scale reading noted This value gives the transmittancy of the sample in per cent To convert transmittancy into optical density the value of $2 - \log R$ is obtained, where R is the scale reading or the table on p 520 may be used Users of this type of instrument appear to prefer the calibration curve method of obtaining results rather than calculation based upon optical density The Hellige instrument is also available in a model ("Clinical") precalibrated for a variety of clinical procedures which are described in an accompanying handbook

The Evelyn photoelectric colorimeter (Fig. 131) is likewise a single photocell photoelectric filter photometer with uniform test tubes customarily employed as solution containers and with readings made on 5

¹ Manufacturers of saturfactory photometers include Central Scientific Co. Cheese Rubseon Co. Philadelpha. Klett Manufacturing Co. New York Fisher Scientific Rittaburgh American Instrument Co. Siver Springs, Vol. Coleman Instruments for May wood III. Helipse Inc. Garden City. N. Y. Photovolt Corp. New York Plauch and Lomb Optical Co. Pochester and others. Some of these convernal kewise manufacture fluorimeters. Information concerning various types a valadie usually may be secured from any laboratory supply house.

sensitive galvanometer which is separated from the rest of the instrument

The galvanometer scale is graduated linearly. from 0 to 100, so that readings here are also in terms of per cent transmittancy A diagram illustrating the schematic construction of the Evelyn instrument is shown in Fig. 132 With a suitable filter in place, the reference fluid in a special test tube is placed in the instrument and the light intensity adjusted by resistance control until the meter reads 100 The reference fluid is then replaced by the sample, in a second similar test tube, and the galvanometer reading noted Its value gives the per cent transmittanes of the sample To convert transmittancy into optical density, the value of 2 - log G is obtuned, where G is the galvanometer reading, or the table on p 520 may he used

The light source is a 6-volt bulh, operated from a storage battery to provide constancy of illimination, which is essential with all single

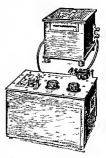


FIG 131 EVELY PHOTO-ELECTRIC COLORIMETER Courtes) Rubicon Co

cell photometers. The filters used with this instrument are particularly satisfactors, from the point of view of narrowness of spectral hand, the selec-

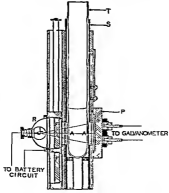


FIG. 132 SCHEMATIC DIAGRAM OF FVELTA PHOTO-LLECTRIC COLORIMFTER.

A, Apertures, F, Filter, L, lamp, R, reflector, S,

shield, T, test tube, P, photocell

tion available 13 shown in Fig 129 The test tubes require a minimum of about 6 ml of solution for a reading, a microcolorimeter, requiring much less fluid, is also available for use with the instrument. The use of test tubes as solution containers or cuvettes has the great advantage that many colorimetric procedures may be carried out partially or wholly in the same tube as will be used for the final reading. The Evelyn test tubes have dimensions such that they correspond roughly to a 2-cm solution thickness, so that the optical density for a particular concentration of substance will be about twice that expected at 1-cm depth, which is the usual basis of reference, and which is used throughout this chapter This means that in the Evelyn instrument, a sensitivity and concentra tion range specified for 1-cm solution depth will correspond to about twice the sensitivity, over the lower half of the specified concentration range only, higher concentrations will prove to be too dark to read accurately This must be considered in interpreting data obtained by this instrument or for use with it

The Bausch and Lomb "Monothromatic" Colorimeter is a single-photocell, direct-reading photoelectric colorimeter, equipped with narrow hand interference filters (See p 525) A 6-volt, 32-c p meandescent lamp operated from a constant voltage transformer, supplies the light for both the scale and the analysis path Light from the lamp passes through condensing lenses, then through a heat-absorbing filter, past a control diaphragm, through an interference filter, and then through the sample solution finally striking a barrier-layer photocell. The current output of the photocell is measured by a sensitive double-suspension galvanometer, the deflection of its mirror (indicated by a floating spot light) being measured upon the translucent scale at the front of the colorimeter

Advantages of single-photocell photometers such as the three just described include the simplicity of construction and the fact that readings are made on a direct-reading meter which does not require manual adjustment Thus even relatively unstable colors, such as that obtained in the antimony trichloride reaction for vitamin A, may be read immediately or at successive small time intervals for extrapolation. The major disad vantage of the single-photocell type is the requirement for a stable light source, to eliminate the possibility of error due to a change from the initial light intensity during a measurement. To provide a constant our rent for the light source storage batteries or constant voltage transform ers are used, the former is generally more satisfactory, since the efficiency of constant voltage regulators may depend upon the type of power supply available Another disadvantage of single-photocell photometers is that they usually incorporate meters designed primarily for current measurement and not for photometric purposes with linear graduation and frequently a small total scale length which may make precise reading difficult Meters should be accurately readable to one-third of a scale division, for example if a reading error of 1 per cent or less is expected over one-third of the scale range. I mear graduation of the scale requires the use of logarithms or conversion tables to convert per cent transmittance values into optical density which is a much more satisfactory basis for photometric analysis. Some types of single-cell photometers are equipped with scales reading in terms of optical density as well as or rather than, per cent transmittancy, it is not unreasonable to hope that this practice will be extended.

To overcome some of the disadvantages of single photocell photometers various types of photometers employing two photocells in a balanced

circuit have heen developed, of which the Klett-Summerson instrument (Fig. 133) is an example. The details of construction of this instrument are shown in Fig. 134. In operation, with a suitable filter in place, the reference solution contained in a test tube is placed in the path of light striking one of the two photocells, which are arranged in a potentiometric circuit so that from the other through a null point instrument (low sensitivity galvanometer). With the photometer scale set at zero (corresponding to zero optical



Fig 133 Klett-Summerson Photo-Electric Colorimeter. Couriesy Klett Manufacturing Co

density) the current output from the second photocell is adjusted so that it exactly halances that coming from the photocell which is subject to the light emerging from the solution. This balance is indicated by a zero reading on the galvatometer. The reference solution is then removed and

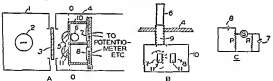


Fig 134 Diagrammatic Details of Klett-Summerson Protoelectric Coloriniter

A Schematic view of the rear half of the instrument in section from above B view across the line 00 in A looking toward the front of the instrument. C Wiring diagram of the photoelectric cell circuit. I Lamp housing. 2 lamp. 3, light filter 4 instrument housing, a compensating lenses 6 test tube 7, working photoelectric cell 8 reference photoelectric cell 9 metal tube (light shield). 10 photoelectric cell compartment housing. II light shits in compartments P 400-ohm potentionater L 100-ohm fixed resistance. G low sensitivity galvanometer.

replaced by the solution under examination. Any light absorption by this solution will throw the two photocells out of electrical balance, balance is then restored by turning the potentiometer dial until the galvanometer again reads zero. The reading on the potentiometer scale at this point is the measure of the light absorption of the solution.

The light source is a 100-watt bulb operated directly from an ordinary power supply, the balanced circuit prevents fluctuations in light intensity from influencing readings. The instrument is designed for use with light filters of relatively narrow spectral transmission, the selection available is similar to that shown in Fig. 129. The test tubes require about 5 ml of solution for a reading, microtubes requiring about 2 ml may also be used. The test tubes may be used for color development as well as for reading and may be centrifuged if necessary. The effective solution depth 1 approximately 1 cm, so that photometric data based on this depth of solution are directly applicable to the instrument. Videls permitting the use of glass cells at other solution depths are also available.

The scale on the Summerson instrument deserves comment, because it is somewhat unusual It is graduated in units which are proportional to optical density, the actual numerical values represent the optical density divided by two and with the decimal point omitted. Thus a scale reading of 250 corresponds to an optical density of 0500, of 100, to 0200 and 40 on. In general, the relation between scale reading R and optical density D is as follows.

$$\frac{1000 \times D}{2} = R$$

Thus the fractional values of optical density (see table on p 520) have been replaced by whole numbers, to facilitate use in photometric calculations. Since the scale readings bear a constant relation to optical density they may be used directly in place of density values in the calculations of photometric analysis.



Fig 135 Perkin Elmer Flame Photometer. Courtesy Perk n Elmer Corp

An interesting recent development in filter photometry is the flame photometry (Figs. 135 and 136). This was designed primarily to facilitate the rapid and accurate analysis of such metallic elements as sodium and potassium—which have intense and characteristic flame spectra. The

^{*}For a review of flame plotometry see Barnes 1 chardson Berry and 1100d Ind-Fing Chem Anal Ed. 17 6d5 (1915) Also Berry Chappell and Barnes Ind Eng Chem-Anal Fd 18 19 (1914)

sample in solution is blown in the form of a fine sprin into a colorless flame, and the intensity of the resulting flame spectrum is determined by the response of a photoelectric cell, as measured on a galvanometer or microammeter. The instrument is calibrated for a particular metal by running a series of standards containing varying concentrations of the metal ion. No ashing or other preliminary preparation of the sample is ordinarily required, sodium and potassium determinations on blood plasma or serum for example require merely appropriate dilution with water. This instrument is known to give excellent results, and has proved to be quite valuable in the particular analyses for which it is fitted.

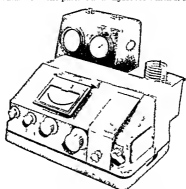


Fig 136 Beckman Model B Spectrophotometer with Flame Photometry Attachment Courtes) Beckman Instruments Inc

Spectrophotometers. In place of using light filters for isolating the narrow spectral region usually required for photometric analysis a device for producing the complete light spectrum and isolating the desired portion may be used Instruments based upon this principle are known as spectrophotometers. In the Beckman photoelectric spectrophotometer (Figs 137 and 138), the spectrum is produced by the use of a quarty prism, a diffraction gruting may also be used, as in the Coleman instrument (Fig 139). The spectral band is focused on a narrow shit mounted in front of the solution being examined and by shifting the band across the plane of the shit the desired spectral region is obtained. Spectrophotometers are usually more expensive than filter photometers the cost increasing with increased sensitivity, spectral range, and narrowness of the spectral region isolated.

³⁹ Manufacturers of spectroplotometers include Beckman Instruments Inc. South Pasadena Calif Coleman Instruments in Maywood III and Central Scientific Co-Cheago Information may also be obtained from the larger laboratory supply houses.

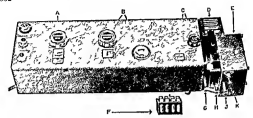


FIG. 137 BECKMAN PHOTOI LECTRIC QUARTZ SPECTROPHOTOMETER.

A, Wavelength scale B_t built in electronic indicating meter, C_t slits with precision adjustment, D_t light source E_t compartment for two phototubes F_t holder for four 10-mm absorption cells, G_t filter slide, H_t compartment for absorption cells J phototube selector, and K_t switch for checking dark current.

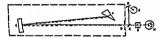


FIG. 138 SCHEMATIC DIACRAM OF BECKMAN

A light B C and E mirrors D shit F prism G envette H phototube

Courtery Cary and Beekman J Opt Soc Am 31 682



FIG 139 COLFMAN MODEL 64 JUNIOR SPECTROPHOTOMETER. Courtery Coleman Instruments Inc

For routine analytical purposes particularly in the visible region of the spectrum spectrophotometers have the advantage over filter photometers of greater convenience and flexibility in choice of wavelength, this is off set to a certain extent by the increased cost as compared to filter photom eters of equal or greater accuracy, and by the increased technical skill required in operation and maintenance. The wavelength setting is an extra adjustment connected with each analysis, variation or alteration is obviously more likely to occur than if a stable glass filter of suitable optical characteristics is used for controlling the spectral range Errors due to maccurate was elength setting may be minimized in an analysis if results are obtained in terms of a standard prepared and read under the same conditions as for the unknown but in any event frequent checking of the accuracy of the wavelength setting is important. The major use of spectrophotometers in photometric analysis at the present time is in connection with substances whose light absorption is in the ultraviolet or infrared regions of the spectrum since no other type of instrument can be used for this particular purpose, and for the establishment of the complete spectral characteristics of a color (i.e., the absorption spectrum)

The Beckman quartz spectrophotometer is especially adapted to meas urements in the ultraviolet and has had wide application in the study of vitamin A, sterols hormones, etc. An infrared model; is also available

Choice of Photometers The choice of photometers from the many types commercially available is largely a question of the requirements of the individual lahoratory For routine analytical purposes hased upon established procedures, any good type of filter photometer will prove satisfactory Instruments equipped for the use of test tubes as solution containers are preferable to other types, because of the convenience and low cost of this type of cuvette, in some laboratories however measurements may have to be made at varying depths of solution and an instrument capable of heing used with different sizes of containers will be required If a single-photocell type instrument is desired, the stability on the laboratory current should be tested before use or facilities for maintaining a storage battery must be available. Filter photometers differ considerably in the width of spectral band transmitted by the filters supplied with the instrument, in general the narrower the band the more satisfactory will be the photometer agreement with Beer's law cannot be expected with filters transmitting a wide range of wavelengths. For analytical purposes a spectrophotometer covering only the visible and neighboring portions of the spectrum is little better than a good filter photometer, except for the convenience of wavelength selection. For investigational purposes the spectrophotometer should be usable in the ultraviolet and near infrared regions as well as in the visible region, an ideal combination would include such a spectrophotometer for investigational purposes and a good filter photometer for analytical purpo es

Turbidimetry and Nephelometry. The light transmittance of a fluid is influenced not only by the amount of light absorbing material present in solution but also by the presence of light scattering or light-obstructing material such as insoluble substances in suspension. Quantitative analysis of substances in suspension based upon this principle is

known as turbidimetry or, less commonly, nephclometry. These two terms are substantially equivalent, although nephclometry is usually considered to include the use of the intensity of scattered light (i.e., light at night angles to the incident beam) as a measure of turbidity, as well as methods based on transmittance measurements. Of the two principles mentioned (light-scattering and transmittance) the latter is more commonly used

It is to be noted that in nephelometric determinations it is not the intensity of the light beam itself that is measured but that of the light scattered at right angles (see Fig. 140). Hence for low concentrations this



FIG 140 KLETT NEPHFLOMETER

The instrument is especially constructed for nephelometric work providing for parallel light which illuminates the cups at right angles. The cups are made of clear glass tubing with black bottoms Courtery Kitch Bhaufactung Co.

method is said to be more sensitive, because a dark background represents zero concentration, whereas there is full illumination at zero concentration in turbidimetric measurements. The chemical principles for producing turbid systems are the same in both turbidimetry and nephelometry and the same precautions must be taken in their preparation and use

Turbidimetric measurements may be carried out by the same procedure and instruments used for the measurement of substances in solution, i.e., by comparison against a series of standards, by dilution, by varying the dight of solution, or by direct measurement of the hight transmitance Transmittance measurement, particularly when used in instruments equipped with photoelectric cells, is the most sensitive and satisfactory. The relationship between the amount of a substance in suspension and the turbidity or transmittance of the fluid is much more empirical than for substances in solution, depending as it does not only on the amount of

material present but also on the size and shape of the suspended particles, their relative opacity or transparency, the relation between particle size and the wavelength of light used, and the uniformity with which a given turbidity may be reproduced. For a particular procedure, bowever, it may be found that over a limited range of concentration the turbidity or transmittance is directly proportional to concentration, and thus resembles optical density or extinction for substances in solution. In such an instance, turbidity measurement is carried out and results calculated in the same manner as for light absorption. In other cases, results must be obtained from a calibration curve constructed from known standards; it is even more important here than for substances in solution that the conditions prevailing at the time the standard turbidities were obtained he reproduced as closely as possible in an analysis.

If the substance in suspension is colorless in a colorless solvent, it may appear at first glance that the choice of wavelength or filter for photometric measurement is immaterial. This is not so, because of the influence of particle size on the scattering of light of different wavelengths ("Tyndall effect"). In general, light of shorter wavelength (at the blue end of the spectrum) is relatively more highly scattered than light of longer wavelength (red end of the spectrum), therefore the change in transmittancy for unit change in turbidity will be greater with short wavelengths than with long wavelengths Thus the sensitivity of the procedure, or the relationship between scale readings and turbidity, may be considerably influenced by the wavelength employed. Other considerations may of course enter into the choice of wavelength; occasionally the interference of extraneous colored material in solution may be minimized by selection

of a wavelength which is not absorbed by such material.

Turbidimetrie measurements are employed not only for analytical purposes but also for the evaluation of partiele size, and for determining the approximate number of plant or animal cells (yeast, bacteria, etc.) present in a fluid; this latter has had wide application, particularly in the field of microbiological assay. Turbidimetrie estimation of red blood cell count has not as yet heen successfully achieved. The use of turbidimetric methods in analytical chemistry has not been great, largely because of the difficulty of achieving a reproducible turbidity for a given concentration. Use has sometimes heen made of a protective colloid to stabilize suspensions and promote uniformity; unfortunately, protective colloids usually distort the relationship between turbidity and concentration in such a way as to require a calibration curve which is highly empirical, and their use cannot be said to have solved the problems of turbidimetric analysis.

Fluorimetry. Certain substances are eapable of absorbing light at one wavelength and radiating a portion of this absorbed light at some other wavelength. This phenomenon is known as fluorescence and may be used for analytical purposes, the intensity of fluorescence serving as a measure of concentration Instruments designed for this purpose are known as fluorimeters. The fluorescence of the unknown may be compared visually against a standard or series of standards, or the intensity of the fluorescent light may be measured directly, using a light-sensitive device such as a

photoelectric cell. Fluorimeters utilizing the latter principle are illustrated in Figs. 141—143

The exeiting light is usually in the ultraviolet portion of the spectrum generally between 300 and 400 mm, and the radiated light is usually in the

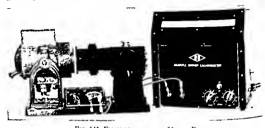


Fig 141 FLUOROPHOTOMETER MODEL B Courtesy Pfalts & Bauer Inc

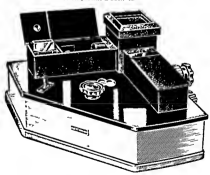


FIG 142 KLETT FLLORIMETER. Courtesy Libra Manufacturing Co

region, but the applications of fluorescence are not limited to these tions. Since the intensity of fluorescence is determined not only by factors which affect light absorption (concentration depth of soliwavelength) but also by the intensity of the exeiting light, measures must be made at a uniform or controlled incident light intensity.

may be obtained from the instructur or from the manual usually supplied by the manufacturer of the instrument? Is it a filter photometer or a spectro-photometer? Note particularly the type of scale used, and whether it is graduated in terms of per cent transmittancy, optical density, or units proportional to optical density. Pface some distilled water in a photometer curetter and insert the cuvette in the instrument. Select any particular filter of wavelength setting, turn on the light, and adjust the photometer to its initial setting, which will represent either zero optical density or 100 per cent transmittancy—which is used? Now remove the cuvette from the light path. Does the adjustment change? Replace the cuvette Does the reading return to its initial value? If it does not, allow the light in burn for about 5 minutes, reset to zero (or 100) and repeat the test of stability. Is there any increase in stability after the light has been on a short while? Why is it necessary for photometer readings to be based on a constant initial setting?

2 Reer's Lau Obtain some defibrinated or oxalated blood Dilute 1 ml of this to 100 ml in a volumetric flask with dilute aimmonia solution (4 ml of concentrated aimmonium hydroxide per litter of water). Mix well by inversion Prepare a series of known dilutions of this "stock standard" as follows into test tubes place 0, 10, 20, 30, etc., up to 100 ml portions of the stock solution, and add sufficient dilute aimmonia solution to each tube to make the final volume 10 ml in each case. Vilx Using light of 520 mg wavelength, adjust the photometer to zero optical density (or 100 per cent transmittance) with the contents of the first tube, which is a "reagent blank". Determine the photometer readings for each of the other solutions, cheking the initial setting with the blank solution once or twice during the series of readings, if any change has occurred reset the photometer hefore continuing with the readings.

using? What must be done in a photometric analysis if (a) the procedure does not obey Beer's law, or (b) a reading falls outside the range of application of Reer's law.

- 3. Relation between Transmittancy and Wavelength. Repeat Exp. 2, but use a wavelength in (a) the blue portion of the spectrum, and (b) the red portion. Plot the results graphically for each wavelength used, and compare with the results of Exp. 2. How may the wavelength chosen influence the sensitivity of a photometric procedure? What is the indication on your graphs of the sensitivity of this particular princedure? Suppose hemoglobin were a contaminant of a solution, instead of the substance being measured; at what wavelength would its presence cause the least error?
- 4. Relation between Transmittancy and Depth of Solution. Repeat Exp. 2, but make measurements at some other depth of solution, if such facilities are available. If not, compute the theoretical reading for each concentration at a depth of solution twice that used in Exp. 2. Plot the data graphically and compare with the previous results. What is the effect of solution depth on the transmittancy value for a given concentration? How may this be used to increase the sensitivity of a photometric procedure?
- 5. Correcting for a Blank Color. Set up the aliquots of 1 100 hemoglobin stock solution used in Exp. 2 (up to 9 ml.) but before diluting to 10 ml. with dilute ammonia solution, add to each tube 0.5 ml of a dilute ammonia solution to which a few drops of blood have been added, sufficient to give a notice-able pink color. Thus each solution now contains a relatively known amount of hemoglobin, plus a constant blank color. Adjust the photometer to its initial settling with the blank tube, and read the other solutions relative to this, as before. Plot the data as described for Exp. 2. Does the presence of a constant blank color influence the photometric calibration, if the photometer is set to its initial reading with the blank solution? Since optical density is additive—i.e., the total optical density of a solution is the sum of the densities due to the various light-absorbing substances present—what other way can you suscess to correctine for the density of a blank?
- 6. Effect of Time of Standing on Transmittancy. Obtain some defibrinated blood, and dilute 0.1 ml. to 25 ml. with 0.1 N hydrochloric acid. Mix hy inversion, transfer a portion to a photometer cuvette, and read immediately, the photometer having been previously set to zero optical density at 520 mm with the dilute acid alone. Repeat the reading on this same sample at suitable time intervals, say every five minutes, for one hour or until the reading becomes constant. Be sure to check the initial setting of the photometer hefore each reading. Many of the colors produced in common photometric procedures show a similar hebation with respect to time of standing. What does this experiment show concerning the desirability of time control in photometric analysis?
- 7. Calibration of Test Tubes Used as Gunettes, Many types of photometers are designed to use interchangeable test tubes in uniform dimensions as solution containers or cuvettes. If the tubes are truly interchangeable, they should all give the same reading for a given colored solution, and they may be tested on this basis. Prepare a stable colored solution of such intensity that it shows by trial a reading about in the middle of the photometer scale.

A dilute solution of hemoglobin, as described in previous experiments, may be used, an equally satisfactory test solution may be made by diluting ordinary india ink with water in a suitable intensity. Place portions of this test solution in each of the test tubes, and read them in the photometer, using water for the initial setting and checking this setting at suitable intervals. Select all those tubes which give identical or nearly identical readings, and discard the remainder Sometimes two lots of tubes may be obtained, the readings for each lot centering with the desired accuracy around a particular reading, but a different reading in each case. Each lot may then be used independently, but they should be marked so that the lots will not be mixed it is also well to mark the tubes so that they will always be placed in the photometer in the same position. Test tubes calibrated in this way are as accurate for photometric purposes as rectangular cuvettes, and much more covenient A tube should be discarded when it becomes scratched, or when checking shows it to have lost its interchangeability.

GENERAL PROCEDURES IN BLOOD ANALYSIS

Drawing Blood for Analysis. Except when analytical methods per mit the use of a few drops of blood (drawn from the finger up, toe, or ear lobe), samples of blood are usually obtained by venipuncture

Procedure Draw a tourniquet (of soft, firm rubber tubing or a strip of bandage) tightly about the arm of the patient a couple of inches above the elbow liave the patient ciench his fist firmly Wash the skin surface about the most prominent vein on the inner surface of the elbow (usually the mediao basilic) with 70 per cent alcohol, allow to dry, hold the vein immobile with the thumb, and into the rein insert a sharp, sterile hypodermic needle (No 18, an inch and a half long) which is attached to a dry sterile syrloge of suitable capacity The needle should penetrate the vein from the side aod at an angle of about 50° with the surface of the arm, the bevel or opening of the needle being kept upward or to the side As soon as blood is seen to eoter the syringe, retract the plunger slowly until the desired amount of blood bas entered the syringe Before removing the needle from the vein, looseo the tourniquet, have the patient unciench his fist, and on the skin at the point of entrance of the needle hold in place a small pad of folded gauze moistened with 70 per cent alcohol Withdraw the needle, detach it from the syringe, and into a suitable container eject the blood from the syringe (not too vigorously, which might cause hemolysis) Pressure on the gauze pad for a few minutes will effectively prevent bleeding from the skin puncture

The use of a syringe is not essential, the needle alone may be used. The blood is allowed to flow from the free end of the needle into a suitable container until the desired amount has been obtained. The needle is then withdrawn as described above. For special precautions to be used in drawing blood where the maintenance of physiological gas tensions is important, as in the determination of carbon dioxide content, etc.

Preparation of Whole Blood and of Plasma. When whole blood or plasma is desired for analysis the blood must be treated with anti-coagulant before clotting commences. The most convenient way to do this to have containers for the blood already prepared with sufficient anti-coagulant in the form of a thin dried film over the inside surface (see be-

low) The thin film promotes quick solubility and mixing with the added blood Test tubes or small wide-mouth bottles of approximately 1-ounce capacity may be used as containers

Procedure Transfer the blood, as quickly as possible after drawing, from the syringe to a container which has sufficient anticoagulant in the form of a thin dried film to prevent clotting of the blood. Mix by rotation gently but thoroughly to dissolve and distribute the anticoagulant. The blood is now ready for use To obtain plasma, centrifuge and remove the supernatant plasma with a rubber-bulb pipet.

Blood specimens are best taken in the morning before breakfast, to minimize the influence of food ingestion. Analyses are preferably made as soon as possible after the blood is drawn, during any interval between drawing and analysis, the blood should be kept cold and well stoppered to minimize evaporation. For blood sugar analyses in particular, the protein-free filtrate should be prepared as soon as possible, to minimize loss of sugar by glycolysis. Protein free filtrates will keep better than whole blood or plasma, particularly if kept cold and in the presence of a drop of toluene as preservativo.

Anticoagulants. The most commonly used anticoagulant is neutral potassium oxalate, of which from 1 to 2 mg are required per ml of blood Only an amount of anticoagulant sufficient for the quantity of blood to be received should be employed, excessive amounts of anticoagulant may interfere with some analyses, cause bemolysis, or produce an abnormal distribution of water and electrolytes between cells and plasma

Procedure To prepare containers with sufficient potassium oxalate for 6 to 0 ml of blood, prepare a stock solution of 10 per cent neutral potassium oxalate and pipet 0 1 ml of this into each container Rotate to produce maximal spreading, then place in an incubator (or oven at 100° C) to dry The oxalate should form a thin dry film on the sides of the container For smaller quantities of blood, use half as much stock solution 11 is good practice to have two sets of containers, suitably labeled, one for 3 to 5 ml of blood, and one for 6 to 10 ml of blood

Other anticoagulants, and the amounts required per ml of blood include sodium extrate (5 mg), lithium or sodium oxidate (1 to 2 mg), sodium fluoride (10 mg), and heparin (0 2 mg) Containers may be prepared with the proper amounts of these anticoagulants in a manner similar to that described above for potassium oxidate. Of the various anticoagulants, heparin is by far the most satisfictory and should be more widely used. Sodium fluoride acts as a preservative and has the advan-

tage of inhibiting glycolytic decomposition of blood sugar, but interfers with certain methods. A special mixture (Heller and Paul)¹⁴ of ammonian salate (3 parts) and potassum oxalate (2 parts) has the advantage of causing no change in red cell volume, and hence is useful for hematoerit determinations and for methods involving the measurement of specific gravity of whole blood or plasma. It cannot be used in ordinary blood analytical procedures because of the presence of ammonia.

Preparation of Blood Serum for Analysis. In the collection of blood for serum, care must be taken to avoid trauma and hemolysis

Procedure When serum rather than whole blood or plasma is desired for analysis, place the freshly drawn blood directly into a small test tube without anticoagulant Allow to clot at room temperature and then chill thoroughly in the refrigerator Centrifuse down the clot and remove the supernatant serum with a rubher bulb ploet

If a centrifuge is not available, the blood may be allowed to clot with the tube in a slanting position, then after chilling overright in an upright position the serum may be poured off from the side of the tube opposite the slanting clot

Measurement of Blood. Because of its physical characteristics and the presence of suspended red cells, whole blood is much more difficult to measure exactly than ordinary fluids. Serious errors have been traced to faulty measurement of blood. Before measurement, the sample must be thoroughly mixed to ensure uniform distribution of cells and plasma.

Procedure To mlk whole blood without trapping all bubbles, if the sample is in a test tube use a small footed stirring rod which is raised and lowered in the sample a sufficient number of times to ensure complete mixing. If the sample is in a wide flat bottle, place the bottle firmly on the table top and vigorously trace a 1 foot circle with the bottle flat against the table top, for at least a dozen times or until the blood is uniformly mixed. Measure out the portion of blood immediately after mixing, and repeat the mixing procedure before such now.

and there is no visible film of blood remaining behind. When the blood has drained completely, blow out the last drop into the receiver. 25

Ordinary pipets may he used for blood measurement, with slight error

due to the fact that they are usually calibrated for the delivery of water rather than blood. Pipets calihrated "to contain" ("TC"), such as are commonly used in microanalyses, are not subject to this error, since they are designed for delivery of the blood into a second fluid, the resulting mixture then being used to rinse out any blood remaining in the pipet. Folin proposed the use of the Ostwald type pipet (Fig. 144), which has less surface per unit volume than the ordinary pipet, and no sbarp shoulders to impede drainage, and these have found favor in many laboratories. Where the amount of blood available is limited and it is desired to use as much as possible, a pipet graduated to the tip is useful (Fig. 145). In using this type of pipet, the greatest error is at the tip, particularly when the tip is imperfect, and if possible the measurement should exclude this portion (i.e., for a 5-ml. portion, measure from the 6-ml. mark to the 1-ml. mark). In general, the most accurate method of measurement is by eareful drainage between two accurately established graduation marks. Volumetric pipets constructed on this principle are not generally available; ordinary graduated pipets should be used in this manner whenever possible.



Fig 145 DILUTING PIPET. Courtesy, Fohn and Wu J Biol Chem. 38, 81 (1919)

PREPARATION OF THE PROTEIN-FREE BLOOD FILTRATE

Method of Folin and Wu.16 Principle. The proteins of whole blood, plasma, or scrum are removed by precipitation with tungstic acid (formed by the interaction of sodium tungstate and sulfuric acid) and filtration. The filtrate is suitable for the determination of the following: nonprotein nitrogen, urea, uric acid, creatine and creatinine, sugar, amino acids, and chlorides Sufficient filtrate for one or two determinations is provided by 2 ml. of blood; for all the determinations, about 10 ml. of blood are needed.

Procedure.¹¹ Transfer a measured quantity of blood to a flask having a capacity at least 15 times that of the volume taken. For each volume (A ml.) of blood taken, add from a buret exactly 7 volumes ($7 \times A$ ml.) of water and

¹⁴ This is for transfer pipets which are calibrated for blowout delivery. Such pipets are commonly marked by the manufacturer with an etched ring around the pipet hear or at the top If the pipet is calibrated for drawings delivery, allow to drain for 1 minute with the tip of the pipet to deliver the wall of the receiver 1 or graduated pipets, allow the blood to run out either to the desired graduation mark, or to the tip if the pipet is graduated to the tip.

Polin and Wu J. Biol Chem. 35, 81 (1919).
 Reagents Required. Sodium tungstate 10 Per Cent Solution. Dissolve 100 g of reagents.

mix. Add I volume (A ml.) of 10 per cent sodium tungstate solution, and mu Flnally add slowly and with shaking I volume (A ml.) of two-thirds normal sulfuric acid. Stopper the flask and shake It. Only a few bubbles should form as a result of this shaking If all the proteins have been precipitated. Let stand for 10 minutes. The color of the mixture should change from red to dark brown. If this change in color does not occur, the coagulation is inconplete, usually because too much oxalate is present. In such an emergency the sample may be saved by adding 10 per cent sulfuric acid, drop by drop with

shaking, until there is no foaming and until the dark brown color has set in
Four the mixture on a dry folded filter large enough to hold it all. Cour
the funnel with a watch glass to minimize evaporation. Collect the filtrate
in a clean dry container. If the first few drops of filtrate are not absolutely
clear, return this portion to the funnel and replace the receiver with a fresh
one. Allow to filter until as much filtrate as possible has been obtained

For plasma or serum the procedure is similar except that 8 volumes of water and only 16 volume of both tungstate and acid are used.

The following modifications of the Fohn-Wu precipitation employ fewer solutions and yield more filtrate.

Hoden's Modification. Add directly to the blood from a buret, 8 volumes of N/12 sulfuric acid. Laking and darkening occur rapidly. Add 1 volumes of 10 per cent sodium tungstate, shake well, and filter, as above.

Von Si₂ke ond Haukins' Modification. Add directly to the blood, 9 rolumes of a mixture of 8 parts N/12 sulfuric acid and one part 10 per cent sodium tungstate. Shake and filter, as above The precipitating solution should be clear and not more than two weeks old.

Biood filtrates prepared by any of the above methods represent a 1 10 dilution of the sample; that is, 1 ml of filtrate corresponds to 0 1 ml of onignal material Since any error in measurement of the added respective will produce a corresponding error in calculations, which are based upon the assumption of an exact 1.10 dilution, all measurements must be made carefully, preferably by the use of calibrated upons to represent the control of the control o

The protein-free filtrates are not and enough to prevent bacteral decomposition If the filtrates are to be kept more than about 12 hour a few drops of toluene or xylene should be added For optimum preepitation of protein, the filtrate should not be more alkaline than pH 28 H A

drop of 0 04 per eent bromophenol blue added to a few drops of filtrate on a test plate should give a yellow or greenish-yellow color, alkalimity being denoted by a pure blue shade

Other Methods of Deproteinization. Although tungstic acid filtrates prepared as described above are the most satisfactory for general purposes, many other reagents have been used for the deproteinization of blood and other biological fluids. Examples of the use of certain of these (zinc and copper bydrovides, triehloroacetic acid, etc.) will be found sub-sequently in this chapter. In most cases they are employed in particular analytical techniques in which tungstic acid filtrates have been found unsatisfactory for one reason or another. Hiller and Van Slyke²² have reported on the comparative value of a number of common protein precipitants. Benedict and Newton²² recommend the use of tungstomolybdic acid in place of tungstic acid for the preparation of blood filtrates, not only for general purposes but also specifically for the deter mination of blood ergothioneine, for which tungstic acid filtrates are unsatisfactory.

DETERMINATION OF NONPROTEIN NITROGEN

Introduction. The nonprotein nitrogen (NPN) of the blood is a collective concept and includes the introgen from all of the nonprotein introgenous constituents of blood which are found in a protein-free filtrate, such as urea, uric acid, creatine and creatinine amino acids glutathione, and many others in small amount, some of which are of unknown nature. Of these various substances, the compound urea contributes by far the largest share to the total, urea introgen representing ordinarily about 45 per cent of the blood NPN. The nonprotein introgen content of blood is usually determined by various micro modifications of the standard Kjeldahl method for the determination of total introgen (see Chapter 31).

1 Method of Folin and Wu ¹⁴ Principle Nitrogen is determined in a portion of the protein free blood filtrate by a micro-Lyeldahl method using a sulfune and phosphoric acid mixture for the digestion, the ammonia formed being determined color methodily after direct nesslengation of the digestion mixture.

Procedure ¹³ Transfer 5 ml of blood filtrate to a large test tube (pyrex) 200 mm × 25 mm, graduated at 35 ml and 50 ml The test tube should either be dry or rinsed with alcohol to reduce the danger of bumping. Add 1 ml of diluted acid mixture and a quartz pebble Boil sigorously over a

Hiller and Van Slyke J Biol Chem 53 253 (1922)
 Benedict and Newton J Biol Chem 33 35" (1929)

¹¹ Benedict and Newton J Biol Chem 33 35 (1929) ¹² Polin and Wu J Biol Chem 33 S1 (1919) For a gasometric method see Van Slyke

J Biol Chem 71 235 (1971)

3 Reagents Required District Acta Warture Vlade by diluting regular acid mixture with an equal volume of water To prepare regular acid mixture add 300 ml of 85 per cent phosphoric acid reagent grade to 50 ml of a 5 per cent copper sulfate solution Add 100 ml of reagent-grade concentrated sulfurne acid and mix heep well stoppered to prevent absorption of ammonis from the air.

Vesiler a Colution "See Appendix."

Sundard Ammonium Sulfiet Soliciton Dissolve exactly 0.23; g of reagent-grade ammon um sulfate in water tras sfer quantitatively to a 1 liter volumetric flask with triusings all a few divergo of concentrated sulfure need make up to volume with water and mix This

Add 2 drops of acetate buffer solution and either 1 ml of urease solution (pre pared the same day) or a piece of urease paper Insert a cork and then either let stand at room temperature for 25 minutes or immerse for 10 minutes in 700 ml of water, having an initial temperature of about 45° C Longer diges tion does no harm if urease paper is used the tuhe must be shaken occasion ally during the digestion period Cool the tube if warm and add an antibump ing tube, 2 drops of antifoaming oli mixture, and 2 ml of saturated borst solution Connect at once with the delivery tube and a test tube recent

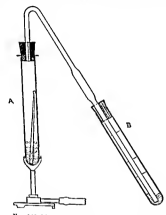


Fig. 146 Microdistillation Apparatus for Urea DETERMINATION Note antibumping tube inside Tube A (Folin and

Svedberg)

graduated at 25 ml as shown in Fig. 146. The receiver contains i ml. of 012 acid and 1 ml. of water. First the contains i ml. of water. acid and 1 ml of water Fasten the boiling tube in a clamp and start the dis tiliation by applying the fiame of a microburner, preferably surrounded by shield to prevent fluctuation of the flame due to air currents. As soon as the contenta are nearly boiling, reduce the flame partially so that the first minut of boiling is very gentle. Then boil briskly for about three minutes and final another minute with the delivery tube slightly raised from the surface o

solve 0 944 g of dry reagent-grade ammonium sulfate in water transfer with rinsings to I liter volumetric flask add a few drops of concentrated sulfurie and dilute to volume at water and mix This solution expression of concentrated sulfurie and dilute to volume and mix This solution expression. water and mix This solution contains 1 mg of nitrogen in 5 ml and is stable indefinitely.

To prepare the dilute standard used in a stable indefinitely and in the standard to To prepare the dilute standard used in the procedure dilute 5 ml of stock standard used in the procedure dilute 5 ml of stock standard used in the procedure dilute 5 ml of stock standard to 100 ml with water This solution contains 0 1 mg of nitrogen in 10 ml and is prepare fresh daily

williquid in the receiver. To another test tube like the receiver, transfer 10 mil of standard ammonium sulfate solution (containing 0 1 mg of N), and 1 mil of 0 1 N acid Dilute both to a volume of about 20 mil. 3 add 2 5 mil of Nessler isolution, dilute to the mark, mix and make the color comparison, using titles a colorimeter or a photometer. For photometric measurement, prepare a blank tube containing 1 mil of 8 1 N acid, water to about 20 mil. add 2 5 mil of Nessler solution, dilute to 25 mil and mix. Set the photometer to zero are density with the blank, and determine the densities of standard and under some solutions as described for the determination of nonprotein nitrogen on p 547 knowns as described for the determination of nonprotein nitrogen on p 547.

CALCULATION For colorimetric measurement

 $\frac{\text{Reading of Standard}}{\text{Reading of Unknown}} \times 0.1 \times \frac{100}{0.5} = \text{mg urea N per 100 ml blood}$

The standard is satisfactor, for blood values between 8 and 40 mg per cent. For higher values repeat the determination with less filtrate plus water.

For photometric measurement

 $\frac{\text{Density of Unknown}}{\text{Density of Standard}} \times 0.1 \times \frac{100}{0.5} = \text{mg urea N per 100 ml blood}$

Under the photometric conditions specified on p 547, the standard has a density of roughly 0300. Up to 60 mg per cent blood urea N may be measured satisfactorily For higher values or with cuveties of greater depth than 1 cm the analysis is carried out on less filtrate plus nater and the calculations corrected accordingly

For separation of ammonia by aeration rather than distillation see the next procedure

Interpretation. Normally, the urea mtrogen of whole blood varies between 10 and 15 mg per 100 ml. On a restricted hospital diet, however values below 20 mg should not be regarded as abnormal. In early nephritis the urea nitrogen may rise to 30 or 40 mg, but in the terminal stages of chronic nephritis and in some cases of acute nephritis marked urea retention may occur. High values may also be found in other conditions associated with damaged renal function, such as mercury bichloride poisoning, double polycystic kidney, intestinal obstruction, prostatic obstruction (in which case the urea N constitutes a valuable guide to surgical risk.), lead poisoning certain infections, cardiae failure, and so on Relatively low figures for urea mtrogen are found in nephrosis (nonhemorthage nephritis with edema), which is probably of metabolic rather than renal origin.

The blood urea concentration has greater significance when interpreted with relation to urea exerction, as in the urea clearance test (Chapter 31)

2 Method of Von Slyke and Cullen ** Principle Whole blood is treated with urease under optimal conditions for conversion of the urea to ammonium carbonate. The mixture is then made alkaline with potassum carbonate and acrated. In the

¹³ In the original procedure of Folm and Svedberg 1 ml of gum ghatti solution (see Appen hit) is a lided at it is point. This is ordinavily not necessary and its use should be a voiled it possil is anneat it decreases color intensity and alters the relationship between citor intensity and concentration. If used it must be added to both atm lard an lunknown in Van Sjk, and Cullen J. Bud Chem. 11, 211 (1914) 24. 117 (1912).

microburner until the characteristic dense fumes begin to fill the tube This will happen in from 3 to 7 minutes, depending on the size of the fiame What the test tube is nearly full of fumes reduce the fiame sharply so that the speed of the boiling is reduced aimost to the vanishing point. Cover the mouth of the test tube with a watch glass. Continue the gentle heating for 2 minuts, counting from the time the test tube became filled with fumes. If the oxidation is not visibly finished at the end of 2 minutes the heating must be continued until the solution is nearly colories. At the end of 2 minutes remote the flame and allow the digestion mixture to cool for 70 to 90 seconds. The add 15 to 25 ml of water. Cool further approximately to room temperature and then fill with water to about I cam under the 35 ml mark. Set aside until the standard is ready, since in this determination it is important that code development be carried out in hoth unknown and standard under as nearly approximate.

A standard suitable for either colorimetric or photometric measurement is prepared as follows transfer 3 ml of a standard solution of ammonium suifate, containing 0 15 mg of nitrogen, to a graduated tube similar to that used for the blood filtrate, add 1 ml of the diluted phosphoric sulfuring acid mixture, to balance the acidity of the unknown, and dilute with water to about 1 cm under the 35 ml mark For photometric measurement a this or blank tube is required, containing 1 ml of diluted acid mixture along made up with water as described for unknown and standard

When all the tubes are ready, nessierize each individually as follows measure out 15 mi of Nessier's solution in a graduated cylinder, set the tube contents in motion by gentle rotatory shaking, and in one continuon motion pour the Nessler solution into the tube Do not wait for the Nessler solution to drain completely from the cylinder, since the exact amount Nessier solution is relatively unimportant, but immediately add water to the tube contents up to the 50 mi mark, Insert a clean rubber stopper, and mix by inversion Proceed Immediately to the nesslerization of the remaining solutions in a similar manner. Allow the solutions to stand 10 minutes after adding the Nessier reagent and mixing, to permit maximum color derelor ment, and read within the next 10 minutes or so Prolonged standing mar lead to the development of a turbidity which renders the color comparison difficult or even worthless 7 Occasionally a turbidity is present before not sierization, due to the action of the acid digestion mixture on the giass of the tube. This turbidity can be removed by centrifuging a portion of the colored solution before color comparison

For colorimetric measurement, match the standard against itsell \$1.20 mm in the usual way, and then compare the unknowns against by standard for photometric measurement, transfer portions of the colorist solutions to sultable containers and determine the densities in the 9hor solutions to sultable containers and determine the densities in the 9hor solutions to sultable containers and the sul

solution is stable indefinitely and contains 0 05 mg of nitrogen per ml Dri the ammonitor sulfate before weighing it out by heating in an oven overnight at 100° C

In all intropens analyses involving the determination of small amounts of amount of the high experience and sessing the state of the high experience and solutions must be known free reagents may be used and solutions must be known tested to prevent the absorption of amounts from the arr Datible with experience and the used to free it from amounts reduct in the presence of a little with the contract of the state of the

tory at by a trap of extrained duties sufficient and protect the distillate from exposure.

2 Calling any of extraining duties sufficient annotate prior to adding the Nessler results has been recommended to good water for that must be prior to adding the Nessler results has been recommended and a few forms of 2 per reen ignum spatis solution (see Appendix) to standard and the all it in old few drops of 2 per reen ignum spatis solution (see Appendix) to standard and the solution is not recommended unless above furly necessary for further ducessoon see footnote 35 p 551.

ometer at 480 to 540 mµ (see below), setting the photometer to zero density (100 per cent transmittance) with the blank

CALCULATION For colorimetric measurement

$$\frac{\mathrm{Reading\ of\ Standard}}{\mathrm{Reading\ of\ Unknown}}\times0\ 15\times\frac{100}{0\ 5}-\mathrm{mg\ of\ NPN\ per\ 100\ ml\ of\ blood}$$

The standard indicated is satisfactory for the range of 15 to 60 mg per cent blood NPN For amounts outside this range repeat the analysis using less or more filtrate as necessary, and correct the calculations accordingly

For pholometric measurement

$$\frac{\text{Density of Unknown}}{\text{Density of Standard}} \times 0.15 \times \frac{100}{0.5} = \text{mg of NPN per 100 ml of blood}$$

The spectrophotometric characteristics of the Nessler reaction with ammonia are illustrated in Fig 240 p 879 For the amounts of nitrogen which are concerned here satisfactory agreement with Beer's law is found with any filter or wavelength setting between 480 and 540 m $_{\rm H}$ the choice depending largely upon the sensitivity desired In a 1 cm cuvette at 480 m $_{\rm H}$ the density of the standard prepared as described above as approximately 0.250 This means that up to about 120 mg per cent blood non-protein nitrogen may be accurately determined under these conditions. For higher values or with deeper cuvettes where the range is proportionately reduced use less filtrate for the analysis and correct the calculations recordingly

Interpretation.*7 The nonprotein introgen content of normal blood ranges from 25 to 35 mg per 100 ml Variations in blood NPN content are ordinarily due largely to variations in urea centent. Increased values for nonprotein introgen are observed in nephritis, more especially in the chronic types and in terminal stages, where they may have prognostic significance, although uremie symptoms may develop without marked elevation of nonprotein mitrogen. The direction of change will import greater information than the magnitude of an isolated determination

In regard to the choice of determining either the nonprotein or the urea N as a clinical test of renal function, the advantages seem to be in favor of the latter, for the following reasons urea covers a relatively wider ringe of variation in disease, it is a single chemical compound rather than a mixture of partly undetermined composition, it is simple to determine clinically, especially if direct nesslerization of the Folm-Wu filtrate is employed. It is of interest, however, that the urea fraction of the non-protein introgen, which usually rises in renal disease, shows subnormal values in cases of celampsia, suggesting an increase in the "rest N" fraction which may contain toxic split-products of protein metabolism

2 Method of Koch and McMeekin "Principle This method differs from the preceding one in that sulfuric acid and hydrogen perovide are used in the digestion

²⁷ For review see Folin Physiol Rers 2, 460 (1922) also Peters and Van Slyke's book (see Bibliography at end of chapter)

¹⁰ hoch and velockin Am Chem See 46 2006 (1921) A stronger and direction mixture committee the said a said a said by Rose J Bool Chem 64 23 (1924) A saturated and unitary committee the said as said by Rose J Bool Chem 15 411 (1925). There is some evidence that both by drogen personde and percoloric said may yield inconstant results possibly it rough our diston of a portion of the ammonia although the proceeding appear to be sufficiently reliable for routine chinella purposes. Selenum is said to be satisfactory as an oxidation catalyst and appears to give good results (see Campbell and Hann J Bool Chem 119 (1937)).

Procedure " Transfer 5 ml of the 1 10 protein-free filtrate to a 200 X 25 mm pyrex test tube, add 1 ml of 1 l sulfuric acid and a small quartz pebble and heat over a microburner to evaporate off the water When charring be gins and white fumes begin to appear in the tube, reduce the size of the flame or raise the tube so that the tip of the flame lust touches the hottom of the tube Continue heating until no further darkening occurs Remore the flame, allow the tube contents to cool for about 1 minute. and then add I drop of 30 per cent hydrogen peroxide, allowing it to drop directly into the solution Replace the flame and heat again to boiling if the solution is not decolorized repeat the addition of the hydrogen peroxide Finally bod gently for 5 minutes Cool, transfer to a 50-ml volumetric flask with about 35 ml of water, and set aside until the standard is ready Test tubes gradu ated at 35 and 50 ml as described for the previous procedure may be used bett Instead of volumetric flasks To prepare the standard, transfer 3 ml of stand ard ammonlum sulfate solution, containing 0 15 mg of nitrogen, to a 50 ml volumetric flask, add 1 ml of 1 1 sulfuric acid, and dilute to about 35 ml with water For photometric measurement a blank is prepared in a third flash by diluting 1 mi of 1 1 sulfurle acid to about 35 ml with water When ready, to each flask add 12 mi of modified Nessier reagent from \$

When ready, to each Hask and 12 mi of modified Nesser reagent prograduated cyiloder, swirling the contents of the flask just before adding the Nessler reagent to promote quick and uniform mixing. Dilute immediately with water to the 50 ml mark, stopper, and mix by inversion Allow to stand 10 minutes before reading. Read in the colorimeter or photometricative as described for the previous method, using the same calculations

Interpretation. See previous method

Other Methods. As has been indicated, most of the modification in the micro-Kjeldahl determination of blood nonprotein nitrogen by direct nesslerization involve changes in the digestion mixturo Alternat procedures for the determination of the ammonia formed include gas metric estimation (Van Slyke, loc cit), aeration of the ammonia from the alkalinized digest as described for the determination of urea (see net section), or steam distillation of the ammonia (see Chapter 31) In bot acration and steam distillation the recovered ammonia is absorbed in acr and may be estimated by titration or by nesslerization, the advantage nesslerization after separation of the ammonia from interfering mater is that crystal-clear colored solutions are invariably obtained A stea distillation device such as that illustrated in Fig 241, p 881, or 1 equivalent, is convenient almost automatic in operation, and requires b a few minutes for each sample Steam distillation prior to estimation ammonia is recommended for all precise micro-Kjeldahl analyses f very small amounts of nitrogen (ammonia) the microdiffusion methof Conway (see pp 668 and 886) has given excellent results

³³ Reagents Required Sulfurse Acad 1 1 With stirring carefully add 50 ml of conferrated sulfure acid to 50 ml of water Cool and keep well stoppered to prevent absorpt of samonia from the sur

²⁰ Per Cent Hydrogen Perorade Only the highest grade low nitrogen reagent (Merck ker a are satisfactory) may be used This reagent is extremely corrosive to the slant is the dispensed with care preferably by the use of a rubber bulb pipet. Keep in Ingerator when not in use. Modified Vester Solution See Appendix.

Standard Ammonsum Sulfate Solution See previous method "

DETERMINATION OF LIREA

Introduction. Practically all of the methods in use at the present time for the determination of the urer content of blood are based upon meubation with preparations of the enzyme urease, 20 whereby urea pres ent is converted into ammonium carbonate. The ammonia formed may then be determined directly by colorimetric methods, or separated by either aeration or distillation and then determined either colorimetrically or titrimetrically. The carbon dioxide produced by decomposition of am monium earbonate may also be measured gasometrically 31 Several methods which do not involve the use of urease have also been described 3" The choice of procedure among the many available appears to be largely a question of the facilities and requirements of the individual laboratory

1 Method of Folin and Stedberg 32 Principle The ammonia produced by the action of urease on the protein free blood filtrate is distilled off and determined colorimetrically by reaction with Nessler's reagent. For a discussion of the accuracy of this method see Gentzkon (J Biol Chem , 143, 540 (1942))

Procedure 4 Transfer 5 ml of tungstic acid blood filtrate to a pyrex test tube of 30 ml capacity (tubes which previously contained Nessler solution should be rinsed with concentrated nitric acid and then with water before use)

¹⁸ Marshall J Biol Chem 15 487 (1913) 11 Van Sly Le J Biol Chem 73 695 (1927)

Ormaby J Biol Chem 146 590 (1942) Barker ibid 152 453 (1944) Archibald ibid

^{157 50&}quot; (1945) See also method of Leiboff and Kahn (p 555)

[&]quot; Folin and Svedberg J Biol Chem 88 77 (1930)

^{**} Reagents Required Acetate Buffer Solution Dissolve 15 g of crystallized sodium acetate in a 100 ml volumetric flash by the help of 50 to 75 ml of water Add 1 ml of glacial acetic acid dilute to volume and mix

Urease Solution Transfer 0.5 g of jack bean meal to a clean 50-ml flash add 20 ml of 30 per cent (by volume) alcohol Shake for 10 minutes and filter or centrifuge This extract should always be prepared on the day it is to be used because on standing even in an icebox it will develop ammonia and will yield too high results. One should therefore not use more extract or a stronger extract than is really necessary. Noch (J. Lab. Clin. Ved. 11 "76" (1926)) obtains a stable and active urease preparation by making a 75 per cent gly cerol extract of jack bean meal

Urease Paper Transfer to a clean 200-ml flask 30 g of tack bean meal and 100 ml of dilute alcohol (30 ml of 95 per cent alcohol diluted to 100 ml) Add 1 ml of the buffer mix ture described above Stopper tightly and shake vigorously for at least five minutes and then shake less hard for about 10 minutes. Filter or preferably centrifuge half an hour in 15-ml tubes the mouths of which have been covered with tinfoil Transfer the extract to a porcelain dish and at once take it up on strips of rather heavy filter paper Schleicher and Sel ull No 597 and hang tlese up to dry over two threads about 15 cm apart While dry ing the papers should not be exposed to air currents for blasts of air seem to destroy the enz; me so long as water is present. As soon as the paper strips are thoroughly dry cut them up into pieces about 1 em by 25 cm and preserve in wide-mouth bottles. These urease papers will retain their activity for many months and even for years. The urease becomes fixed in the paper and it is only by shaking the solution several times during the digestion

that one secures adequate contact and quantitative hydroly as of the urea histomping Tibe. As illustrated in Fig. 14. Made preferally from pyrex glass 2 mm in diameter at open end May be obtained from 1sider Securities Co. New York histomping Oil Virture To one volume of crude fuel oil add about 10 volumes of

toluene Saturated Boraz Solution Dissolve about 40 g of reagent-grade sodium tetraporate (borax) in I liter of boiling water and allow to cool to room temperature If unsaturated ad I more borax and again heat

^{0.1 \} Acid Either by droel loric or sulfurie may be used and it need not be standardized Standard Ammonium S Unite Solition Prepare a stock standard solution as follows Dia

as described in the original procedure of Van Slyke and Cullen. The aeration procedure may also be applied to the Folin-Wu filtrate after urease treatment, the liberated ammonia being absorbed in dilute acid and determined colorimetrically. The ammonia aerated from urease-treated whole blood may also be determined colorimetrically instead of by titration.

3. Method of Karr: Principle. By the action of urease, the area in a protein-free filtrate is converted to ammonium carbonate which is nesslenzed in the presence of a proteintie colloid. The interference of peptones and ammo acids (Folin and Wu) is regarded as so slight and uniform as not to influence the clinical value of the results.

Procedure.¹² Transfer 5 ml, of Folin-Wu blood filtrate to a test tuhe, and in a similar tube place 5 ml. of standard urea solution, containing 0.075 mg. of nitrogen. To each tube add 0.5 ml. of buffer solution and 5 drops of urease solution. Place in a bath at 50° C. for 15 minutes. Transfer the contents of each tube, with rinsings, to separate test tubes** graduated at 22.5 and 25 ml. and dilute with water to the 22.5 ml. mark. Add 3 drops of gum ghatti solution, followed by Nessler solution to the 25-ml. mark. Alix, allow to stand for 10 minutes, and read within the next 20 minutes, using either a colorimeter or photometer. For photometric measurement, follow the conditions specified on p. 547 for the determination of nonprotein nitrogen, setting the photometer to zero density with a blank prepared by treating 5 ml of water with buffer, urease, etc., exactly as deseribed above for a blood filtrate.

CALCULATION For colorimetric measurement

Residing of Standard Reading of Unknown
$$\times 0.075 \times \frac{100}{0.5} = \text{mg}$$
 ures N per 100 ml blood

For photometric measurement

Density of Unknown Density of Standard
$$\times 0.075 \times \frac{100}{0.5} = \text{mg}$$
 urea \rightarrow per 100 ml blood

Interpretation. See under method of Folin and Syedberg above.

o Karr J Lob Clin Med. 7, 3 (1924) The use of guin ghatti as a stabilizing colloid by Folin (J Biol Chem., 81, 231 (1929) in his sugar method (p. 575) suggested its application contacts. In Comput. J. May 2012, 1923.

in the urea method to Looney [J. Biol Chem., 88, 189 (1930)

"Reagents Required Standard Urea Solution Davslove 0.2215 g of pure dry urea in water and dilute to 500 ml in a volumetric flack. Add a little chloroform or tolures as a preservative This solution contains 0.3 mg of urea introopen per ml To prepare the worther standard, dilute 5 ml of stock standard to 100 ml with water in a volumetric flack and mix. Prepare fresh daily This solution contains 0.075 mg. of urea intropen in 5 ml

Buffer Solution Dissolve 20 g of crystalline sodium acetate in water add 2 2 ml of 10 per cent aceta acid dilute to 100 ml with water and mix Add a little toluene or chloroform as preservative

Urease Solution Place 15 g, of jack bean meal (obtainable from the Arlington Chemical Co Yonkers, NY), about 2g of Permutit' (see Appenday) 16 ml of 95 per cent alcohol, and 84 ml of water in a 200-ml flank. Shake more or less continuously for about 16 minutes. Four onto a filter and allow to filter overnight in the refrigerator Keep the filtrate in cell transferring a portion to a dropping bottle for daily use Prepare fresh every timed. Gum Obatts Solution See Appendix.

Nester Solution The Koch-McMeekin preparation is recommended (see Appendix) "Separate tubes for nesselection are necessary because of the 'poisoning' action of mercury on urease When this is suspected in the conversion tubes they should be cleaned with strong intre acid

4 Method of Leiboff and Kahn " Principle. The ures in the Folin-Wu blood filtrate is converted to ammonia by acid hydrolysis under pressure and directly nesslerized. The slight conversion of nonurea substances to ammonia is unimportant clinically By the elimination of urease, turbidity after nesslerization is claimed to be avoided even with filtrates high in urca

DETERMINATION OF CREATINING

Introduction. Blood ereatinine is ordinarily determined by reaction in a protein-free filtrate with alkaline picrate to form a red color (the Jaffé reaction), which is then compared with a standard A color reaction of creatinine with dinitrobenzoate has also been described 45 It is well recognized that the Jaffé reaction is by no means specific for creatinine. and that other substances are present in blood (chiefly in the red cells) which contribute to the color, so that results on whole blood filtrates are undoubtedly too high 46 Practically all of the chromogenic material in plasma, bowever, appears to be ereatinine and therefore plasma is preferred to whole blood for analysis Most of the results in the literature have been obtained on whole blood. A specific method for creatinine determination appears to be the measurement by the Jaffe reaction before and after treatment with a hacterial preparation which destroys creatinine 47

Method of Folin and Wu " Principle A portion of the blood filtrate is treated with alkaline picrate solution and the color developed is compared in a colorimeter or photometer with that produced by a known amount of creatinine under the same conditions

Procedure 4 Transfer 10 ml of 1 10 tungstic acid filtrate of whole blood or plasma (preferably the latter) to a small flask or test tube. In a second container, place 5 ml of standard creatinine solution, containing 0 03 mg of creatinine, and add 15 ml of water Add 5 ml of freshly prepared alkaline picrate reagent to the blood filtrate, and 10 mi to the diluted creatinine

⁴⁴ Leiboff and Kahn J Biol Chem 83 347 (1929)

⁴⁵ Benedict and Behre J Biol Chem. 114 515 (1936) Langley and Evans ibid. 115 333

⁴⁶ Hunter and Campbell J Biol Chem 32 195 (1917) Behre and Benedict ibid 52 (1922) 117 415 (1937) Willer and Dubos ibid 121 447 457 (1937) Gaebler and Abbott ibid 123 119 (1938) 47 Miller and Dubos loc cut Allinson J Biol Chem 157 169 (1945)

⁴⁴ Folin and Wu J Biol Chem 38 S1 (1919) See also Peters J Biol Chem 146 179 (1942) Bonenes and Taussky biod 158 S51 (1945)
48 Reagents Required Standard Creatinine Solution Prepare a stock standard by dissolv

ing 1 g of pure dry creatinine in 0 1 N by drochloric acid and diluting to 1 liter with the seid This solution is stable indefinitely and contains 1 mg of creatinina per ml To prepare the working stan lard transfer 3 ml of stock standard containing 3 mg of creatinine to a 500ml volumetric flask add 50 ml of 0 1 N by drochloric acid dilute with water to 500 ml and mix This standard contains 0 03 mg of creatinine in 5 ml and is atable for a week or mora if preserved by the a idition of a few drops of toluene

Alkaline Purate Reagent (a) Prepare a saturated solution of purified picric acid (see Appendix) It is essential that this solution be saturated otherwise serious error may result 110-ml portion titrated with 0 1 \ alkali in the presence of phenolphthalein as indicator should require 5 2 to 5 4 ml of alkali for neutralization (b) To prepare the fresh alkaline picrate reagent transfer 25 (or 50) ml of the saturated picric acid solution to a flask add 5 (or 10) ml of 10 per cent sodium hydroxide solution and mix Use witl in a short time after preparing On standing crystals may form in the solution which do not impair its effectiveness but rei der measurement difficult

556

standard Mix and allow to stand for 15 minutes for complete color devel opment Read in the colorimeter or photometer within the next 15 minutes

For colorimetric measurement, first match the standard against itself carefully, and then compare with the unknowns in the usual way For photometric measurement, transfer the solutions to suitable containers and determine the densities in the photometer at 520 mm (see Fig. 148) Set the photometer to zero density with water alone Determine the density of a hlank prepared by treating 10 mi of water with 5 mi of the alkaline picrate reagent, and subtract this value from the observed densities of standard and unknown to obtain their true values

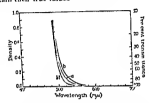


FIG. 148 ABSORPTION SPECTRA OF COLORED SOLUTIONS OBTAINED IN FOLIN WE BLOOD CREATINING METHOD

Alkaline picrate slope (II) alkaline picrate plus 0.03 mg creatinine (a) plus 0.06 mg creatinine (b) Solution depth 1 cm

CALCULATION For colorsmelese me sourement

Reading of Standard \times mg. creatinine in standard $\times \frac{100}{1} \times \frac{10}{30}$

= mg creatinine per 100 ml 1 lood or plasma

In connection with this calculation it is to be noted that the standard is made up to twice the volume of the unknown so that a volume correction factor (154a) is introduced. The standard specified corresponds to a blood creatinine of 15 mg per cent and is satisfactory for the range of 1 to 2 mg per cent only because of the presence of the high blank color of the alkaline picrate and the significant deviation from Beer's law (see p 557) slown by this color reaction. It is good practice therefore to prepare several standards at different concentration levels to provide for values outside the indicated range. If a high creatinine should be encountered without several stand ar is ready the determination may be saved by diluting the unknown with an appropriate amount of the alkaline pierate solution which I as first been diluted with two volumes of water to preserve equality of pieric acid an l alkali concentration 1t is better lowever to repeat the analysis if possible using less filtrate plus water to 10 ml correcting the calculations accordingly

For photometric measurement values up to about 5 mg per cent blood creatinine may be calculated as follows

Density of Unknown \times mg ereatinine in stan lar $1 \times \frac{100}{1} \times \frac{15}{10}$

= mg_ereatinine per 100 ml 1 lood or plasma

For higher values repeat the analysis with 5 ml of filtrate plus 5 ml of water and rultiply the results by 2 4t 520 mµ and ma 1-em envette, the density of the stand and described corrected for the blank is approximately 0 050.

This calculation is an approximation only, suitable for routine clinical purposes but not for precise work since it is based on Beer's law, and the creatinine color does not follow Beer's law exactly at any concentration ordinarily encountered. For more exact work a calibration curve relating observed densities to concentration must be prepared and results read from this curve. To obtain such a curve prepare a standard creatinine solution which is twice as strong as that used in the procedure by trans ferring 6 ml of the stock 0 I per cent creatinine solution to a 500-ml volumetric flask adding 50 ml of 0 1 N acid and diluting to the mark with water. This solution con tains 0.06 mg of creatinine in 5 ml. Transfer the following amounts of this solution (preferably in duplicate) to small flasks 00 50 100 150 and 200 ml Add sufficient water to each flask to make the final volume 20 ml, and then add 10 ml of alkaling picrate reagent to each Mix allow to stand 15 minutes and determine the densities in the photometer under the conditions described in the text. The first flask is a blank subtract its average value from the other average values to obtain their true densities The amounts of creatinine in the various flasks are 0.06 0.12 0.18 and 0.24 mg respectively corresponding under the conditions of

analysis to blood creatinine contents of 3 6 9 and 12 mg per cent. Plot the true densities against the equivalent mg per cent concentrations on cross-section paper and draw a smooth curve to include all the points A curve similar to that shown in Fig. 149 should be obtuined. From such a curve the concentration of an unknown may be read off if its density has been established.

In using such a curve, it is important to remember that it is valid only if the analysis of an unknown is carried out under conditions similar to those prevailing at the time the curve was established. In general, the calibration for one photometer is not applicable to another instrument even of the same make and changing the wavelength setting or filter will influence the curve. The curve should be checked when a new lot of saturated pieces well as prepared although correcting for the blank as indicated will ordinarily take care of slight variations in the color of the alkaline

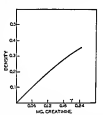


FIG 149 TYPICAL CALIBRATION CUNVE FOR BLOOD CREATINAT DETERMINATION AT 520 Mµ 1 CM SOLUTION DEFIN

picrate reagent. Time control and temperature control are important, the curve at 30° C is not the same as at 20° All of these precautions are necessary for obtaining accurate results

Interpretation. Creatume is the least variable introgenous constituent of the blood, in which it exists to the extent of 1 to 2 mg per 100 ml of whole blood, the average value for plasma is nearer 1 mg per cent. In early nephritis, values of from 2 to 4 mg are noted, and in chronic hemorrhagic nephritis with uremia, 4 to 35 mg. Creatume is more readily excreted by the kidners than urea or uric acid, and an increase of creatumine to 4 or 5 mg or over per 100 ml is exidence of marked impairment of kidner function. Such high creatumine values in chronic hemoriem.

³⁶ Peters (loc cit) recommends a pieric acid solution containing 11.75 g per liter rather than a saturated solution to minimize differences in solubility of pieric acid at summer and winter temperatures

rhagic nephritis indicate an unfavorable prognosis, although high values may obtain in acute cases over long periods.

DETERMINATION OF CREATINE

Introduction. The creatine of blood is ordinarily determined by the Folin-Wu procedure of heating the protein-free filtrate with acid, which converts creatine to creatinine The total creatinine present is then determined by the procedure used for blood creatinine. The value for preformed creatinine, as established by separate analysis, is subtracted from the total creatinine value, to give the amount of creatine, expressed as creatinine. There is no satisfactory method for the direct determination of blood creatine. The specificity of the method based on acid treatment and the Jaffé reaction for creatinine is open to question Increased specificity is undoubtedly obtained by the use of the creatinine-destroying bacteria of Miller and Dubos;11 few applications of this principle to knowledge of blood creatine have as yet been made.

Procedure, Transfer 5 mi. of a 1:10 tungstlc acid filtrate of whole blood to a test tube graduated at 25 ml. Add 1 ml. of normal hydrochloric acid. Cover the mouth of the test tube with tin foll and heat in the autoclave to 130° C. for 20 minutes or to 155° C. for 10 minutes. Cool, Add 5 ml. of freshly prepared alkaline picrate solution (as used for blood creatinine determination) and let stand for 8 to 10 minutes, then dilute to 25 mi. At the same time, prepare a standard creatinine solution by adding to 10 ml. of creatinine solution, containing 0.06 mg. of creatinine, in a 50-ml. volumetric flask, 2 ml. of normal acid, and 10 ml. of the alkaline picrate reagent, and after 10 minutes, diluting to 50 ml. Read in the colorimeter or photometer as described for the determination of blood creatinine (p. 556). The blank for photometric measurement is prepared by treating 10 ml. of water in a 50-ml. volumetric flask with 2 ml. of acid and 10 ml. of alkaline picrate reagent and diluting to 50 ml. with water.

In the case of uremic bloods containing large amounts of creatinine, 1, 2, or 3 ml. of blood filtrate, plus water enough to make approximately 5 mi., are substituted for the 5 ml. of filtrate, and calculations corrected accordingly.

CALCULATION For colorimetric measurement

Reading of Standard Reading of Unknown $\times 0.06 \times \frac{100}{0.5} \times \frac{25}{50} = \text{mg}$ "total creatinine" per 100 ml blood

Subtract the preformed creatmine content, as determined by separate analysis (p 555), from the "total creatinine," to obtain the creatine content, expressed as creatinine The same precautions as were described for the determination of blood creatinine, concerning deviation from Beer's law and the use of several standards if necessary, must be observed

For photometric measurement

Density of Unknown Density of Standard $\times 0.06 \times \frac{100}{0.5} \times \frac{25}{50} = \text{mg}$ "total creatinine" per 100 ml blood

Determine creatine by subtracting preformed creatinine value as described above

al Allinson J Biol Chem. 157, 169 (1945) For application of the bacterial method to the determination of creatine in tissues, see Miller Allinson, and Baker J Biol Chem., 130, 333 (1939), Borsook and Dubnoff. 55ad. 132, 559 (1940)

As with the determination of creatinme the calculation given is only an approximation because of deviation from Beer s law and for precise results a calibration curve similar to that described on p 557 should be constructed and used. The calibration curve prepared for the preformed blood creatinine determination cannot be used here because of the presence of acid and the different proportion of alkaline picrate used. In preparing this curve, remember that only 5 ml of filtrate are used for analysis in stead of 10 ml as in the creatinine determination so that a given amount of creatinine in the standard corresponds to twice as high a total creatinine content per 100 ml of blood as in the calibration for preformed creatinine. Thus the standard described containing 0.06 mg of creatinine in a final volume of 50 ml is equivalent in an analysis to a total blood creatinine of 6mg per cent with other amounts in proportion

Interpretation Blood creatine content as determined by this and other methods ranges hetween 3 to 7 mg per 100 ml of whole blood. The creatine of blood is found almost entirely in the red cells, variations in cell count therefore presumably influencing results. Except in the case of infants and pregnant women, the amount of creatine in plasma is quite small, Allinson (loc ctl) reports values ranging from 0.4 to 0.8 mg per cent. So No significant variations have as yet been associated with pathological conditions, so the determination has little elimical value at the present time Increases have been noted after severe muscular injury and following experimental removal of the kidnoys.

DETERMINATION OF URIC ACID

Introduction. Blood uric acid is ordinarily determined by colori metric procedures hased upon the reaction between uric acid and certain complex phosphotungstic acid reagents (or their equivalent), usually in the presence of cyanide 22 to form a hlue color Procedures utilizing the reducing action of uric acid on ferricy anide have also been described 64 The determination of blood une acid has been the subject of much criticism and study. It appears probable that a portion of the blood une acid is lost during precipitation of the proteins, up to now, there appears to be no simple way to avoid this 55 The use of the various color reagents on a blood filtrate directly, without prehminary separation of interfering material, is felt by many investigators to be unsound because of both the nonspecificity of the color reaction and the possible presence of substances which inhibit color development with uric acid itself. A combination of these effects (one tending to increase color, the other decreasing it) is felt by some to be responsible for the apparently satisfactory results obtained in some procedures. Interfering material appears to be present largely in the red cells of blood, the use of plasma or serum for analysis is therefore gaining ground and is to be preferred. Increased specificity is also obtained by separating the urie acid from interfering material prior to nnalysis Color development before and after treatment with preparations of the enzyme uricase, which oxidizes urie neid, has also been proposed

¹² See also Tierney and Peters J Clin Invest 22, 595 (1913)

³⁷ For a procedure not involving the use of cyanide see hern and Stransky Biochem Z.,

³⁴ See Hulger and Jol ns J Biol Chem 140 42" (1941)

[&]quot; See lowever hern and Stransks (loc cut)

as a basis of specificity 56 Of the three procedures described below, two are hased on separation of interfering material before color development The third, the method of Brown, is a direct method which appears to be as satisfactory as any that have been described, and simpler than most

Method of Folin (Isolation Procedure) " Principle Unic acid is precipitated as silver urate, directly from the blood filtrate. The uric acid is set free by means of acid chloride solution and determined colorimetrically or photometrically after the addition of phosphotungstic acid, which gives a blue solution

Procedure * Transfer 5 ml of the blood filtrates to a centrifuge tube Add 2 ml of the acid aliver solution Centrifuge at once All the uric acid, down

44 Blauch and Koch J Biol Chem 130 443 (1939) For a description of a uricase procedure applied to the determination of une acid in urine see Chapter 31 | Urine Quantita

tive Analysis
U Folin J Biol Chem 101 111 (1933) 106 311 (1934) The Folin direct method 15 similar to the procedure described here and requires the same reagents except that the precipitation with acid silver solution is omitted A 5-ml portion of the blood filtrate is treated directly with the cyanide-urea solution uric acid reagent ete exactly as described in the text Calculations and directions for photometric and colorimetric measurement are the same

16 Reagents Pequired Acid Silver Solution To 5 ml of 85 per cent lactic acid add 100 ml of water and 5 g. of NarCOs and boil Dissolve 20 g. of silver nitrate in about 700 ml of water add the partly neutralized factic acid solution and dilute to I liter After a few days exposura to sunlight and filtering this reagent keeps fairly well When used only occasion-

ally however it should be filtered before using

Standard Une Acid Solution The solution made as follows will keep for at least fire years Weigh out on a watch glass exactly i g of ure acid and transfer it to a liter volumetric flask hy means of a not too small dry funnel. Tay the funnel co as to transfer nearly the whole of the uric acid to the flask. Transfer 0 6 g of lithium earbonate to a 250-ml Florence flask, add 150 ml of water chake about five minutes until dissolved Soma insoluhie material remains and it is usually best to filter. Heat the solution or filtrate to 60° C Also warm the liter flask under running warm water Pour the warm lithium earbonate solution into the liter flask incidentally washing into it the traces of uric acid which ad hered to the watch glass and funnel. Shake so as to dissolve the urie acid promptle. A little additional warming under hot tap water is permissible. The lithium carbonate solution is not always perfectly clear even when filtered and one should not mustake this little surb d to for undesolved uric acid and keep warming and shaking too long. In five minutes all of the uric acid should be dissolved Shake the flask under cold running water without undue delay Add 20 ml of 40 per cent formalin and half fill the flask with dis illed water Finally add from a pipet rather slowly and with shaking 25 ml of normal sulfuric acid Dilute to volume mix thoroughly and transfer to a clean tightly stoppered bottle This stock solution containing I mg of une seid per ml should be kept away from light

To prepare the working standard dilute i ml. of the stock solution with water only to 200 ml It behaves exactly like a lithium carbonate solution of uric and and keeps perfectly

for many days. (o ml. = 0 02 mg, urse acid)

Urea-Cyanide Solution (Poisonous) Transfer 75 g. of Merek a Blue Label and um cyanide to a 2-liter beaker add 700 ml of water and stir until the cyanide is completely dissolved Add 300 g. of ures and stir Then add 4 to 5 g. of calcium oxide and stir for about 10 min ntes Filter at once if necessary for immediate use but preferably not until the next day To the filtrate add about 2 g of powdered fithium oxalate shake occasionally for 10 to 15 minutes and filter

Une Acid Reagent Transfer 100 g of sodium tungstate (free from molybdate of footnote 59 below) to a .00 ml. Florence flask. Mix 32 to 33 ml. of 80 per cent pho-phoric acid with 150 ml of water Pour the resulting solution on to the ting-tate and mix. Add a few pebbles and boil very gently over a microburner for I hour Loss of liquid during the boiling is prevented by using as a condenser a funnel lolding a 200-ml flask filled with cold water At the end of the boil ng period decolorize with a little bromine water boil off the excess

bromine cool and dilute to 500 ml

If the reagent so obtained is not perfect (in other words if it gives a blank with Merck a urea-cyamide or with urea-cyanide plus tyro-une) add 3 to 5 g, of sodium tungstate (but to the lost trace, will now be in the preclutote Decant the supernatont solution as completely as possible, and add I ml of o 10 per cent solution of sodium chloride in 0 1 N hydrochloric acid Stir thoroughly with a fine glass rod, add 4 ml of water, and stir again Centrifuge Pour the supernotont solution as completely as possible into a test tube graduated at the 25 ml mark. Remove the last drop by touching the lip of the centrifuge tube to the side of the graduated test tube Prepare two standards by placing 3 ml (plus 2 ml of water) and 5 ml respectively of standard uric acid solution, contolning 0 02 mg of uric acid in 5 ml . In separate test tubes, and for photometric measurement prepore a blank tube containing 5 ml of water only To each tube odd 10 ml of urea cyanide solution (Poisonous' from a buret) and mix well by lateral shaking Add 4 ml of the uric ocid reagent to each tube, mix well by lateral shoking, and note the time Let stand for 20 minutes, dilute to the 25 ml mark with water, stopper, and mix by inversion Read in the usual way within the next half hour in the colorimeter or photometer. For photometric measurement, determine the densities of 420 mu, setting the photometer to zero density with the blank

CALCULATION For colorsmetric measurement

Reading of Standard \times mg urise acid in standard $\times \frac{100}{0.5}$

= mg une acid per 100 ml blood or plasmn

Read the unknown against the standard which most closely matches it on inspection and substitute the proper value for the une acid content of the standard (0 012 or 0 020 as the case may be) in the calculation formula above. The stronger standard corresponds to a blood with 4 mg per cent une acid up to 8 mg per cent may be read satisfactorily. For higher values instead of using the entire supernatant from the acid extraction of the precipitated silver urate as described pipet out 2 5 ml at this point add 2 5 ml of water and continue with the addition of urea-cyanide color development etc. as described multiplying the final results by 2.

For pholometric measurement

 $\frac{\text{Density of Unknown}}{\text{Density of Standard}} \times \text{mg uric acid io standard} \times \frac{100}{0.5}$

= mg uric acid per i00 ml blood or plasma

Either standard may be used therefore only one is necessary. At 420 m μ and in a 1-cm cuvette the density of the stronger standard corresponding to 4 mg per cent blood urie and is approximately 0.500 Up to 8 mg per cent may be accurately read under these conditions. For higher values or with deeper cuvettes use half of the supernation fluid from the acid extraction as described above under colorimetric measurement and multiply the results by 2

no more) and boil for another 10 to 15 minutes then cool and decolorize as before Tl c add tion of a little extra tungstate and the abort second boiling can also be made without first testing the reagent for a blank

^{**} The tungsite acid deprotenmention preceding estimation of ure said by this method must be made with sodium tungstate entirely free from molybdate A product meeting this specification is made by the Vallinckrodt Chemical Co. The absence of molybdate elouid be confirmed on eith new batch of sodium tungstate by applying the following xanitate test (Folin and Trimble J Biol Chem. 60 173 (1924)). Dissolve I g of sod um tungstate in 5 to 10 ml of water. The solution of solid be alkalme Add about 0.2 g of solid potassium ethyl annit ate (Fatnian or prepare as directed in original paper). Stake until dissolved Add drow we with alkang 20 per cent In Foli annit the curfy tungstate in Frequently and a public to deep thin color forms. The colored compound molybdenium sanithate is soluble in teleprotum.

At 520 mµ and in a 1-cm cuvette the standard described has a density of 0 180 (Fig. 180). Under these conditions measurements are reliable up to 10 mg per cent ure and For higher values or with a deeper cuvette use less filtrate plus water to 5 ml in the acid silver chloride prespitation and correct the calculations accordingly. The color in this procedure does not reach a constant value changing gradually in intensity with time. This change takes place at a bout the same rate in the standard and in the unknowns. In serial analyses it is necessary, therefore, to reread the standard at suitable intervals and to calculate results in terms of the density of the standard at approximately, the time it e density of an unknown is determined.

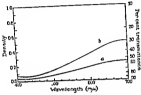


Fig. 150 Absorption Spectra of Colored Solutions Obtained in Newton Blood Uric Acid Method

For standards containing 0.01 mg uric acid (a) and 0.02 mg, uric acid (b) Solution depth 1 cm

Interpretation. Results on human blood by this procedure are closely similar to those obtained by the previous method. Occurrence of turbidity is much less frequent than for other methods. For further interpretation see p. 562

3 Method of Brown ** Principle The tungstic acid filtrate is treated directly with a special une acid reagent in the presence of optimal amounts of evanuations colution. The color developed is compared with that of a unic acid standard treated similarly.

Procedure "Transfer 2 ml of 1 10 tungstic acid filtrate to a test tube of cylinder graduated at 10 ml. In a similar tube place 2 ml of standard uricacld solution containing 0 005 mg of uricacld, and for photometric meas

⁶⁵ Brown J Biol Chem 158 601 (1945)

[&]quot;Peagents Required Softum Cyanude Solution 12 Per Cent Dissolve 12 g of pure sodium cyanide in water in a beaker transfer to a 100-ml cylinder add water to the mark and mis with a strings not Handle this solution carefully as the sectional positions as keep in the refragerator It should be usable for about 2 weeks Bring to room temperature before

transferring a portion to a buret for daily use

Urea Sol aton Disselve of g of urea in sufficient water to make 100 ml This solution
keeps indefinitely at room temperature

Une Acid Reagest Dissolve 100 g of reagent-grade sodium tungstate and 20 g of and 5 drous dissolution by drogen ph spl at ein about 150 ml of water in a flask with the aid of test this 25 ml of concentrated sulfare and with about 25 ml of water and pour the warm solution aboutly and with slashing into the flask. Place a funnel in the moutt of the flask and in the funnel place 200-ml flask filled with the water Heat the matture in the

tube add 2 ml of cyanide solution from a buret, this reagent is highly poisonous and must near be dispensed from a pipet. Nix by lateral shaking, and add 2 ml of urea solution to each tube. Again mlx. Finally add 1 ml of the urle and reagent, mix by lateral shaking, and allow to stand at room temperature for 50 mlnutes. Dilute to the 10 ml mark with water, stopper, and mlx by Inversion. Read in the colorimeter or photometer in the usual way. For photometric measurement, read at 520 mµ, aetting the photometer to zero density with the blank.

CALCULATION For colorimetric measurement

$$\frac{\text{Reading of Standard}}{\text{Reading of Unknown}} \times 0.005 \times \frac{10}{2} \times 100 = \text{mg uric acid per 100 ml blood}$$

The standard described corresponds to a blood une acid content of 2.5 mg per cent For values over 5 mg per cent repeat the analysis using 1 ml of filtrate plus 1 ml of water and multiply the results by 2

For photometric measurement

$$\frac{\text{Density of Unknown}}{\text{Density of Standard}} \times 0.005 \times \frac{10}{2} \times 100 = \text{mg uric acid per 100 ml blood}$$

At 520 m μ and in a 1-cm cuvette the density of the standard (corrected for the blank) is approximately 0 200. Satisfactory agreement with Beer's law is found up to blood concentrations of 6 mg. per cent. For higher values or with deeper cuvettes, use 1 ml of filtrate plus 1 ml of vater for the analysis and multiply the results by 2

Interpretation. By this method human whole blood shows a normal range of 2 2 to 3 5 mg per cent for males and 1 9 to 2 9 mg per cent for females Results are lowered by only about 10 per cent or less when the uricase procedure (see p 559) is applied to establish specificity For further interpretation, see p 562

DETERMINATION OF AMINO ACIDS

Principle The color developed by the reaction between amino acids and β-naphthoquinone-i sulfonic acid in alkaline solution is the basis of this method. Originally proposed by Fohn the method was considerably improved by Danielson " and this procedure is essentially the one described here with heating to develop the color as suggested by Sahyun " and with photometric measurement according to Frame, Russell and Withelmi ".

Procedure 70 Prepare a tungstic acid filtrate of whole blood or plasma at a 1 10 dilution in the usual way (see p 543) Transfer 5 ml of protein free

flask to boiling and boil gently for I hour the funnel arrangement serving as a condenser Cool and transfer to a I I ter volumetric flask with rinsings dilute to il e mark with water and mix

and mix

Ure: Acid Standard Dilute 1 ml of the Folius stock urse acid standard containing 1 mg
of ure acid (see footnote 85) to 400 ml with water and mix This solution contains 0 005 mg
of ure acid in 2 ml and lects well for several days particularly it kept cold.

[&]quot; Danielson J Bul Clem 101 503 (1933)
" Sahyun J Lab Clin Mel 24 548 (1938 1939)

I I I rante Russell and Willelni J Biol Chem 149 255 (1913) See also Russell J Biol Chem 156 467 (1944) for a slightly modified procedure claimed to give better results are Recurrent Amino Leaf Standard Sol Itom A mixed standard containing gly

cine and glutanic acid is recommented for colorimetric measurement because of better match with Hood filtrates Clipciae Sandart D soulce exactly 0.08 g of pure dry glycite in water aid transfer with rinsings to a 500 ml volumetric flask \ \dd 35 ml of \ \ \hat{N} \hat{groups}

Interpretation.40 Normal human whole blood usually contains from 2 to 3 5 mg, of uric acid per 100 ml, by this method. The normal range for plasma is 3 to 5 mg per cent. In early nephritis values of from 3 to 10 mg. may be noted, and in advanced cases values as high as 25 mg, are observed The determination of blood une acid is generally regarded as an unsatisfactory index of renal impairment since its level mny be affected by a chemical destructive process as well as by excretion, which might account for the observed inconsistencies between hyperuricemia and other nonprotein nitrogen retention.

In gout, high uric acid values (4 to 10 mg) are usually found. Determination of une acid is of diagnostic value in gout prior to the stage of tophi formation, particularly since normal values, in the absence of salicylate or dietary therapy, would ordinarily exclude a positive diagnosis However, uric acid is similarly increased in early nephritis and many eases of gout with high urie acid values also show defective kidney function by other tests The same difficulty is met with in considering the high values (2 to 8 mg) obtained in other arthritic conditions, usually associated with increases in urea also The existence of nephritis in such cases has not been entirely excluded and many typical cases of arthritis show values below 3 mg. Salieylates and atophan (cincophen) tend to reduce the uric acid content of the blood.

The excessive breakdown of nuclear material in leukemia is accompamed by elevated uric acid values. In addition to the numerous conditions resulting in renal damage, such as mercury or lead poisoning, eclampsia, acute infections, mahgnancy, etc , the blood uric acid is also increased in certain eardise conditions which ultimately involve the kidneys.

2. Method of Newton: Principle. Interfering material in the blood filtrate is removed by the acid silver chloride precipitation method of Benedict and Behre. The uric acid remaining is determined colorimetrically by reaction at room temperature in the presence of eyanide with a special arsenotungstate reagent,

Procedure.4 The blood proteins are precipitated with the molybdotungstic acid reagent of Benedict and Newton rather than by the Folin-Wu tungstic acid method, although this latter filtrate may also be used. Transfer 1 volume of blood to a small flask and add 7 volumes of water to lake. Add I vol-

⁶⁰ See reviews by Folin, Berglund, and Derick J Biol. Chem , 60, 361 (1924), and Rose Physiol Revs. 3, 544 (1923) See also Myers and Muntwyler: Ann. Rev. Biochem. 9, 303

Newton J. Biol Chem. 129, 315 (1937).
 Benedict and Behre J Biol Chem. 92, 161 (1931) For the complete uric acid procedure of these authors, see the eleventh edition of this book.

⁴¹ Resgents Required For Protein Precipitation Tungitomolybdate Solution Boil 10 g of rengent-grade ammonia-free molybdic acid with 50 ml of normal sodium hydroxide for 4 to 5 minutes Filter, and wash the residue on the filter with 150 ml of hot water Cool the filtrate and washings and add to a solution of SO g of sodium tungstate dissolved in 600 ml of water Dilute to I liter and max Q62 h Sulfuric Acid Dilute 620 ml of N sulfuric acid to 1 liter with water, and mix. Both these solutions are stable indefinitely Acid Lithium Chloride Dissolve 7.5 g of lithium chloride in water add 35 ml of concen-

trated hydrochloric acid dilute to I liter with water, and mix Stable indefinitely

reason systematics can ensure to I liter with water, and mix Stable indefinitely.

Siter Natural Solution Dissolve 29 g of reagoni-grade siver intract in water, dilute to

liter, and mix Keep in a brown bottle

Acid Bank Solution (Or photometric measurement only) Dilute 7 ml of concen-

ume of tungstomolybdate solution followed by I volume of 0 62 N sulfuric acid, with shaking Allow to stand for a few minutes, then pour onto a dry filter and collect the filtrate in a small dry flask. For plasma or serum, use 8 volumes of water and 0 5 volume of the tungstomolybdate and acid

Transfer 5 ml of the I 10 filtrate prepared as described to a 15-ml centrifuge tube, add 1 ml of acid lithium chloride, and mix Add 1 ml of silver nitrate solution and shake well Centrifuge at once and pour the supernatant fluided late a test tube, allowing time for complete drainage. Touch the llp of the centrifuge tube to the test tube to obtain the last drop. In a second test tube place 5 ml of the standard uric acid solution, containing 0 02 mg of uric neld, and add 2 ml of water For photometric measurement prepare a third tube containing 5 ml of "acid blank" solution plus 2 ml of water To each tube add from a buret 3 ml of the cyanide-urea reagent (poisonous), mix by lateral shaking, and follow with 1 ml of the diluted lithlum arsenotungstate reagent from a buret (poisonous) Stopper and mix by inversion Allow to stand 10 minutes, and then read in a colorimeter or photometer For photometric measurement, determine the densities at 520 mu, setting the photometer to zero density with the blank

CALCULATION For colorimetric measurement

Reading of Standard $\times 0.02 \times \frac{100}{0.5} = \text{mg}$ uric acid per 100 ml blood or plasma

With the standard at 15 mm readings of the unknown between 10 mm and 30 mm are reliable. For higher values, use less filtrate plus water to 5 ml in the acid silver chloride precipitation and correct the calculations accordingly

For photometric measurement

Density of Unknown $\times 0.02 \times \frac{100}{0.5} = \text{mg}$ uric neid per 100 ml blood or plasma

trated hydrochloric acid to I liter with water and mix

Cyanide Urea Reagent (Foisonous') Dissolve 100 g of urea in 500 ml of water Add 25 g of reagent grade sodium cyanide and stir to dissolve This solution is usable for I month or so Fresh solution must be prepared if color develops in a blank test run as described for photometric measurement in the text

Lithium Arsenolungstate Reagent (Poisonous') Dissolve 100 g of reagent grade sodium tungstate in 500 ml of water and add 140 g of reagent grade arsenic pentoxide Boil under a reflux condenser for 1 hour Remove the condenser and continue boiling until the volume is reduced to about 200 ml. Pour this solution onto 100 g of solid lithium chloride in a beaker Stir until all of the lithium chloride has gone into solution. Chill the contents of the beaker to at least 10° C for 2 hours but no longer Filter off the precipitated lithium arsenotungstate by suction and suck as dry as possible. Excess water may also be removed by pressing the precipitate between filter papers. About 130 g. of material should be obtained. Dissolve in water and dilute to 500 ml. This stock solution keeps indefinitely. Before use a portion is diluted with 4 volumes of water. This diluted reagent is used in the procedure The salt may also be prepared in quantity preserved in the dry state and dissolved in water as needed

Standard Uruc Acid Solution (Benedict) A atock solution is prepared as follows Dissolvo 0 g of disodium hydrogen phosphate and 1 g of sodium dihydrogen phosphate (pure crystalline salts) in about 200 to 300 ml of hot water if not perfectly clear filter Dilute the clear solution to 500 ml with hot water and pour upon exactly 200 mg of pure uric acidsus-pended in a few ml of water in a liter solutinetric flask. Mix until solution is complete. Cool add exactly 1 4 ml of glacial acetic acid dilute to mark and mix Add 5 ml of chloroform to prevent bacterial or mold growth (5 ml - 1 mg of urie acid) The working standard is prepared fresh weekly as follows Transfer 10 ml of stock solution to a 500 ml volumetric flask dilute to 400 ml ad l 3 5 ml of concentrated hydrochloric acid dilute to mark and

mix (5 ml = 0 02 mg of uric acid) 44 A slight op-lescence may be noted due to a trace of colloidal silver of loride but this disappears after the addition of the cyanide solution and does not affect the accuracy of the results.

filtrate to a test tuhe graduated at 15 ml. In a similar tube place 5 ml. of amino acid standard solution, containing 0.03 mg. of amino acid nitrogen, and for photometric measurement prepare a blank tube containing 5 ml. of water. 11 Add I drop of 0.25 per cent alcoholic phenolphthaleln solution to each tube, followed by 0.1 N sodium hydroxide solution drop by drop until a permanent pink color is obtained. Adjust by adding a little water where necessary so that all tubes are at approximately the same volume. To each tube add I mi. of horax solution, mix by tapping, and then add I mi. of freshly prepared naphthoquinone solution. Mix by tapping, and place the tubes immediately in a boiling water bath. Allow to remain 10 minutes, then remove and place in cold water for 5 minutes. To the cooled contents of each tube add 1 mi. of acid-formaldehyde solution, mix immediately, add 1 mi. of 0.1 N andium thiosulfate solution, dilute immediately to the 15-mi, mark with water, and mix by inversion. Allow to stand for 10 to 30 minutes before reading in the colorimeter or photometer. For colorimetric measurement, match the unknown against the standard in the usual way For photometric measurement, transfer the solutions to sultable containers and read in the photometer at 490 mg, using the blank solution for actting the photometer to zero density.

CALCUI ATION For colorimetric measurement-

Reading of Standard $\times 0.03 \times \frac{100}{0.5}$

- mg amino acid nitrogen per 100 ml blood or plasma

If the unknown has an amino acid nitrogen content outside the range of 4 to 8 mg per cent, repeat the determination using smaller portions of filtrate or standard as required, made up to 5 ml with water, and correct the calculations secondingly.

For photometric measurement

Density of Unknown × 0 63 × 100 Density of Standard

- mg amino acid nitrogen per 100 ml blood or plusma

For the spectrophotometric characteristics of the color obtained in this method, see Frame Russell and Wilhelm (foc cit.) At 490 mm and in a 1-cm cuvette the density of the standard is approximately 0.400 at 520 mm, which may be used with slightly less accuracy the density is about 25 per cent less. Up to about 15 mg per cent amino acid nitrogen may therefore be determined accurately under these conditions for higher levels repeat the analysis with a smaller portion of filtrate

Interpretation. The amino acid nitrogen content of whole blood as determined by this method ranges from 5 to 8 mg per cent. A slight inerease is noted after the ingestian of protein foods but no significant change is found in fasting High values have been observed in uremic nephritis although a remarkable constancy is observed in most pathological conditions Increases have, bowever, been noted in leukemia and particularly in acute yellow atrophy of the liver. Insulin markedly reduces the amino acid content of blood Plasma averages around 4 to 5 mg per cent, with serum sbglitly bigher according to MacTady en,72 this increase is due to release of amino acids during clotting. The concentration of amino nitrogen in corpuscles is about twice that of whole blood. Of naturally occurring interfering substances, glutathione is of chief significance, since this substance is present largely in the corpuscles, interference from this source is minimized if plasma (or an "unlaked" blood filtrate⁵⁷) is taken for analysis. Uric acid at a concentration of 1 mg per cent gives color equivalent to 0 1 mg per cent amino acid nitrogen, 4 correction may be necessary on bloods with high unc acid content. The sulfonamides also interfere if present, a free sulfoaamide level of 10 mg per cent being equivalent to 0 8 mg per cent amna acid nitrogen, 60 correction may also be applied here Despite these limitations, the method is considered to be reliable and satisfactory for blood analysis

Other Methods. Other colorimetric methods are not as satisfactory as the one described here Gasometric methods include the classical mitrous acid method of Van Slyke, "which is based upon the measurement of introgen liberated from a amino acids on treatment with introus acid, and the gasometric ninky drin method of Van Slyke and Dillon," in which the carbon diovide liberated from the carbonyl group of a-amino acids on boiling with minhy drin is measured. For description of the ninky drin method as applied to the determination of the amino acid content of urine, see Chapter 31. Microbiological methods for the determination of individual amino acids are described in Chapter 33.

DETERMINATION OF GLUCOSE

Introduction. The majority of methods for the determination of blood glucose are based upon the ability of glucose in hot alkaline solution to reduce certain metallic ions, of which the cupric and ferricy andeions are most commonly used. The extent of reduction is then established by colorimetric, titrimetric, or gasometric methods. Methods involving or

¹³ MacFadyen J Biol Chem 145 387 (1942) The Van Slyke and Petera Qualitative Clinical Chemistry Vol II (Methods) Baltimore Williams and Wilkins Co 1032

⁷⁴ Van Slyke and Dillon Proc Soc Exptl Biol Wed 34 362 (1937) Van Slyke Dillon MacFadyen and Hamilton J Biol Chem 141, 627 (1941) MacFadyen loc cit

including yeast fermentation. 18 while undoubtedly the most specific for glucose, are not used routinely. It has long been known that there are reducing substances other than glurose present in blood, and that these may occur in sufficient amount to increase considerably the "annarent" plucose value The term "saetharoid" has been proposed to designate the non-glucose reducing fraction of blood 26 The various blood sugar methods differ in their specificity for glucose, and therefore tend to give slightly or even significantly different values for both normal and pathological blood in using a method, or in interpreting results obtained by it, particularly in the early literature, it is important to know the relationship between the values obtained by the particular method and the "true glucose" values, as well as the values obtained by other and possibly more specific methods. It is also important to know whether the method employs venous or arterial (capillary) blood, since this may influence the interpretation of results (see p 577) Because of the free diffusibility of glucose between red cells and plasma, distinction between the analysis of whole blood and plasma is relatively unimportant except for methods which include the reduction due to saccharoids, which are found chiefly in the red cells

I Method of Folin and Wu " Principle The protein free blood filtrate is heated with sikaline copper solution using a special tube to prevent reoxidation. The cuprous oxide formed is treated with a phosphomolybdic acid solution, a blue color being obtained which is compared with that of a standard

Procedure 74 Transfer 2 ml of the tungstic acid blood filtrate (or 1 ml plus i ml of water if very high blood sugar values are expected) to a Folin-Wu

307 (1933) Fast ens and Stiff field 137, 21 (1941)) the saccharoid fraction of hormal human flood is accounted for almost entirely by glutathione and glucuronic acid

¹⁷ Folin and Wu J Biol Chem 41 367 (1920) Folin (J Biol Chem 67, 357 (1926) 82 83 (1929)) has critically examined and improved this method to obtain a more accurate measure of true sugar and better proportionality between sugar concentration and color intensity The original Folin Wa method is here described because it continues to have Nide usage especially in hospital laboratories

18 Reagents Required Standard Sugar Solutions Three standard sugar solutions should be on hand (a) a stock solution 1 per cent glucose made up in saturated benzoic acid solu tion (b) a solution containing 2 mg of sugar in 1 ml (20 ml of stock solution diluted to 100 ml with water) (c) solutions containing 0 2 and 0 4 mg of sugar in 2 ml made by dilution of (b) with water The dilute standards are best made up fresh a couple of times a week. Merck a highest purity destrose is satisfactory

were, were a signess purify destrose is satisfactory.

Alkdinic Copper Solution Dissolve 40 g. of pure sinhydrous sodium carbonate in about 400 ml of water and transfer to a liter flask. Add 7 5 g. of tattaric acid and when the latter has dissolved 60 d. 6 g. of crystallized copper sulfate. Wix and make up to a volume of the companion of the co liter If the chemical, used are not pure a sediment of cuprous oxide may form in the course of one or two weeks. If it is should happen remove the clear supernatant reagest with a aiphon or filter through a good quality filter paper. The reagent seems to keep indefinitely To test for the absence of cuprous copper in the solution transfer 2 ml to a test tube and add 2 mi of the molybdate phosphate solution the deep blue color of the copper should almost completely vanish in order to forestall improper use of this reagent attention should be called to the fact that it contains extremely little alkali 2 ml by titration (using the fading of the blue copper tartrate color as indicator) requiring only about 1.4 ml of

Phosphomolybdic Acid Solution To 35 g of molybdic acid and 5 g, of sodium tungstate add 200 ml of 10 per cent sodium hydroxide and 200 ml of water Boil vigorously for 20

¹¹ See Somogni J Biol Chem. 28, 117 (1928) Van Slyke and Hawkins. 35.5 (1929) Holden 3rd, 119.347 (1937) Wintler Science, 99.327 (1944) "Benedict J Biol Chem. 21.14 (1931) According to Fashens (J Biol Chem. 198

sugar tube graduated at 25 ml (Fig. 151) and to other similar tubes add 2 ml of standard sugar solutions containing 0 2 and 0 4 mg respectively of glucose To each tube add 2 ml of the alkaline copper solution. The surface of the mixtures must now have reached the constricted part of the tube. Transfer the tubes to a rapidly bolling water bath and heat for 8 minutes Cool in

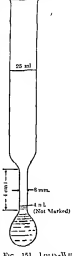
running water without shaking To each tube add 2 ml of phosphomolybdic acid reagent. After about 1 minute dilute to the mark with water and mix. It is essential that adequate attention be given to this mixing because the greater part of the blue color is formed in the bulb of the tube Compare In a colorimeter using the standard which most nearly matches the unknown For photometric measurement, transfer the solutions to suitable containers and determine the densities at 420 mu, setting the photometer to zero density with a blank obtained by treating 2 ml of water with alkaline copper reagent, heating, etc. just as in the analysis of the blood filtrate

CALCULATION For colorimetric measurement use the standord which most closely matches the unknown and calculate as follows

Reading of Standard \times mg glucose in standard $\times \frac{100}{0.2}$

= mg glucose per 100 ml blood

Unless the readings of the standard and unknown are within a few millumeters of each other results obtained by this calculation are somewhat incorrect because the color is not strictly proportional to the concentration of glucose Correction of observed blood sugar volues" may be of importance wheo the values are near the critical levels correspond ing to hypoglycemia and to the threshold range Instead of applying corrections it may be advisable to employ in place of the usual standards, standards containing 0 1 and 93 mg of glucose in 2 oil (corresponding to blood sugar values of 50 and 150 mg per 100 ml respectively) or to dilute to an approximate color match (using the graduated sugar tubes of Rothberg and Lyans) to prior to matching in the colorimeter



151 | LOLIN-WU SUGAR TUBE

For photometric measurement only one standard and a blank are required, and the calculation is as follows

Density of Unknown \times mg glucose in standard $\times \frac{100}{0.2}$

= mg glucose per 100 ml blood

At 420 mm and in a 1-cm cuvette the density of the standard containing 0.4 mg of glucose, corresponding to 200 mg per cent of blood glucose is approximately 0 300 (Fig. 152) Satisfactory agreement with Beer's law is found under these conditions up

to 40 minutes so as to remove nearly the whole of the ammonia present in the molybdie acid Cool dilute to about 350 ml and ad 1 125 ml of concentrated (85 per cent) phosphoric acid Dilute to 500 ml

¹⁰ Over and harr (J. Biol. Chem. 67, 319 (1926)) have published tables and curves correcting for the deviation from Beer a law

⁴² Rothberg and Evans J Biol Chem 33 413 (1923)

to about 400 mg per cent blood glucose. For higher values, or with deeper cuvetteerry out the analysis using less fill rate plus water to 2 ml and correct the calculations accordingly.

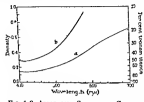


FIG. 152 ABSORPTION SPECTRA OF COLORED SOLUTIONS OBTAINED IN FOLIN W.L. BLOOD SCGAR METHOD For standards containing (a) 0.2 mg. and (b)

For standards containing (a) 0.2 mg and (b 0.4 mg glucose Solution depth 1 cm

Interpretation The normal range of fasting venous blood sugal values by this method is 90 to 120 mg per 100 ml of whole blood Thirange is undoubtedly "enhanced" to the extent of possibly 20 to 30 mg per cent by the effect of non gluose reducing sub-tances (saccharoids) but the saccharoid fraction appears to be relatively constant and therefor its presence does not influence unduly the interpretation of variations il blood sugar level as obtained by this procedure. Nost of the early data if the American literature on blood sugar content were obtained by the method, but it is gradually being superseded by the more specific method described subsequently.

In mild debetes a large of 10 to 2002.

In mild diabetes values of 140 to 300 mg per cent are obtained, and in severe diabetes values up to 1200 mg per cent have been noted. Hyperglycemia is also observed in severe neighbrits paincreatic disease hyperthyroidism, and certain hepatic disorders. Experimentally, the administration of adrenaline and the induction of ether anesthesia leave to elevated blood sugar values. Low blood sugar values are found after the administration of insulin in such hypoendocrine disturbances a Addison's disease, hypoprituatisms cretinism and myxedema, and in the chinical condition known as hyperinsulnism. For carbohydrate tolerances, especially of the condition of the condit

2 Method of Benedict "Principle The protein-free filtrate is heated with a alkaline copper reagent containing tartrate alanine and bisulfite. The reduced copper is determined colormetrically after treatment with phosphomolybdic acid." Talkaline copper reagent used here is practically unaffected by the nonsignar reducing materials present in blood hence the method gives values distinctly lower than with the Folin Wu method and presumally prearr the true glucose content. The reager is also adapted to the determination of the nonsignar reducing value of blood (storaginal papers).

^{*} Benerict J Biol Chem. 92, 141 (1931)

Procedure 12 Place 2 ml of 1 10 protein free filtrote¹³ in a Folin Wu sugar tube, and in a second similar tube place 2 ml of standord glucose solution To each tube add 2 ml of the copper reagent containing bisuifite Mix by lateral shoking, ond place in a vigorously boiling water bath for 6 minutes Cool by placing in cold water for 1 to 2 minutes (avoid shaking at this stage) To eoch tube add 2 ml of color reagent, mix by vigorous lateral shoking, and ofter about 1 minute dilute with water to the 25 ml mark. Mix the content thoroughly by repeated inversion, allow to stand at least 10 minutes, ond read in the colorimeter or photometer within the next 30 minutes or so For colorimetric measurement, compare against the standord in the usual way For photometric measurement, transfer portions of the solutions to suitable containers and determine the densities in the photometer under the same conditions as described for the Folin Wu method, p 569

CALCULATION Same as for Folin Wil method

This procedure gives better proportionality between color and concentration than the Folin Wu method. A standard glucose solution containing 0.2 mg of glucose equivalent to 100 mg per cent blood sugar is therefore suitable to colorimetric comparison for values up to 200 mg per cent blood sugar. For higher levels use a stronger standard or less filtrate plus water to 2 ml. In photometric measurement agreement with Beer's law is excellent up to 500 mg per cent blood sugar at 420 mg and in a 1-cm cuvette. For higher values or with a deeper cuvette earry out the onalysis using less filtrate plus water to 2 ml. and correct the calculations accordingly.

Interpretation Normal values on whole blood by this method range from 70 to 100 mg per cent blood glucose. The interpretation of variation from the normal range is similar to that already presented in connection with the Folin-Wu method.

3 Somogy: Shaffer-Hartmann Method "Principle Hemolyzed blood is deproteinized with zine hydroxide giving a filtrate containing practically no reduced substances other than sigar The sugar is estimated by indometric thration of reduced.

¹⁸ Reagents Required Copper Reagent Dissolve 15 g of anhidrous sodium carbonate 3 g of alanne and 2 g of Rochelle salt in about 250 m of distilled water Dissolve 3 g of crystalline copper sulfate in about 100 ml of distilled water and add it is solution with stirring to the carbonate-alanne-tartrate solution Dilute to 500 ml and mix Keep in a cool lace. This reagent will keep ready mixed from 4 to 6 weeks. If after several weeks there may be a slight growth of modd in the solution it is be removed at any time by pouring the solution through a loose plug of absorbent cotton in a funnel leaving the efficiency of the reagent unaffected.

I Per Cent Sodium Bisi lite Solution Kept in a 100-ml dropping bottle The solution al ould be prepared fresh about once a month

Copper Reagent Containing Buruffix Measure into a cylinder a volume of the copper reagent which will be used up in one or two days and add 1 drop of the bruffits addition for each ml of copper reagent or 1 ml of bisulfits for each 20 ml of reagent Mix Do not use after the second day.

Color Reagent To 180 g of pure molybdie seed and 75 g of anhydrous Na; CO; ma large Frienmey er flask add 500 ml of water in small portions while shaking fleat to boiling litter Wash residue on filter with hot water until filtrate plus wasl ings equal 600 ml Add 300 ml of 85 per cent HiPO; cool and didute to 1 liter Standard Cli cost Solution See footnote 78 p 568.

¹¹ Benedict uses a tungstomoly bdate filtrate (Benedict J Biol Chem 92 135 (1931)) A

¹ oim Nu l'Itrate may also be used "Somogry J Biol Chem 86 655 (1930) 78 509 (1976) Urea creatinine and creatine may also be determined in the rine filtrate but only traces of uric acid are found Lower but significant values for nonprotein aircrogen may be obtained Q. Biol Chem 87 370 (1970) For an improved proceedure suitable for colorimetric as well as titrimetric determination see Somogry J Biol Chem 189 61 69 (1945)

copper Precipitation of the protein with copper salts is just as satisfactory in the case of whole blood and is better for plasma or serum as

Procedure (a) DEPROTEINIZATION Take I volume of blood with 7 volumes of water Add I volume of 10 per cent solution of ZnSO4711:0 Mix Add with continuous shaking I volume of 0 5 N NaOII Stopper the flask, shake well and filter after a few minutes through dry filter paper For accurate work measure the blood with an Ostwald pipet calibrated "to contain" and rinse with the laking water For serum or plasma dilute with 8 volumes of water and add 0 5 volume of each of the reagents Another procedure for whole blood is to add 8 volumes of an acid zinc solution (125 g of ZnSO, 711,0 and 125 ml of 0 25 N 11,80, with water to make 1 liter) and 1 volume of 0 75 N NaO11 Shake vigorously and filter after a few minutes 50 ml of the acid zinc solution should require 67 to 68 ml of 0 75 N NaOll to give \$ permanent pink with phenolphthaleln Determine sugar as below For microtechnique introduce into a test tube or 25 mi Erlenmeyer flask 5 8 mi of water, add 0 2 ml of blood from an accurate capillary pipet, rinsing several times with the laking water Mix Add 1 mi of 1 8 per cent ZnSO, 7li,0 Mix Add with continuous shaking 1 ml of 0 1 N NaOli " Stopper, shake well, and filter after a few minutes through a dry thin paper (Schieicher and Schüll No 597, 70 mm) Use 5 ml of filtrate equivalent to 0 125 ml of blood Deter mine sugar by procedure described

(b) DETFRMINATION OF SUGAR. Measure 5 ml of the copper reagent into 2 large test tube (25 × 250 mm) and add 5 ml of the sugar solutions contain ing not less than 0 1 nor more than 2 0 mg of glucose Shake gently to mis, cover the tube with a small funnel, bottle cap, or glass bulb, and keep in a boiling water bath for 15 minutes Cool (avoid shaking) by placing in a shal low dish of water until the temperature falls to 35° to 40° C (not below 30°) Add 1 ml of 5 N II;SO, (or equivalent amount) and see that all Cu;O is promptly dissolved Avoid reoxidation of reduced copper or loss of iodine by too vigorous agitatioo After about 2 minutes titrate with 0 005 N sodium thiosulfate using starch as an indicator toward the end of the titratioo Run a blank on 5 ml of reagent after heating with an equal volume of water

CALCULATION From the blank tetration sultract the titration of the unknown This gives mi of the issulfate required For the glucose equivalent consult the following table which applies to the usual i 10 dilutions of blood. For other cases the actual amount of glucose in the 5 mi of solution used for the determination is obtained by dividing the value in the talle by 200

¹⁸ Somogyr J Biol Chem 90 725 (1931) Lake I volume of blood in 7 volumes of water Add 1 volume of 7 per cent CuSO, 5HO and mrx Then add with continuous shaking 1 vol ume of 10 per cent NajWO 2HrO Stopper the flask. Shake well Filter through dry filter paper after a few minutes For plasma or serum use the same procedure but use 5 per cent copper sulfate and 6 per cent sod um tungstate solut ons

¹⁰ ml of the rine sulfate solution diluted with 60 ml of water and slowly titrated with the NaOH should require 12 to 12 2 ml for a permanent pink with phenolphthalein 17 Copper Reagent Dissolve 12 g of Roci elle salt 20 g, of sod um carbonate (anhy drous) and 25 g of sodium bicarbonate in about 500 ml of water Into this pour with attring 6.5 g of copper sulfate (crystalline) dissolved in about 100 ml of water Dissolve 10 g of potations and de 0.80 sum nod de 08 0g (weighed to eg) of potass um nodate and 18g of potassium nosaiste in bout 200 ml. of water Add to the main solution and diute to I liter

This sulfate For preparation of 0 t N solution see Appendix D lute this every day or two as needed to 0 005 N

³³ Strongly acid sugar solutions must be neutralized to phenol red. For above filtrates of tungatic acid filtrates this is not necessary

Amounts of Glucose Corresponding to Titration Values When 5 ml. 1-10
Blood Filtrate and 5 ml. Copper Reagent (Modified) Are Heated
In Water Bath for 15 Minutes

Milliliters of 0 005 N Thio- sulfate	Tenths of 1 ml of 0 005 N Sodium Thiosulfate									
	0	1	2	3	4	5	6	7	8	9
	Mg Glucose in 100 ml Blood									
0			21	23	26	29	31	34	36	39
1	41	44	46	49	51	53	56	58	61	63
2	65	68	70	72	75	77	80	82	84	86
2 3	89	92	94	97	99	101	103	106	108	110
4 5	113	115	117	119	121	124	126	128	130	132
5	135	137	139	141	143	146	148	150	152	154
	157	159	161	163	165	168	170	172	174	176
0 7 8 9	179	181	183	185	187	190	192	194	196	199
8	201	203	205	207	210	212	214	216	218	221
9	223	225	227	230	232	234	237	239	241	243
10	245	248	250	252	254	256	259	261	263	265
11	267	270	272	274	276	279	281	283	285	288
12	290	292	294	296	299	301	303	305	308	310
13	312	314	316	318	321	323	326	328	330	332
14	334	337	339	341	343	345	347	350	352	354
15	356	359	361	363	365	367	370	372	374	376
16 17	378 400	381	383	386	388	390	392	394	396	398

Interpretation. Normal values by this method range from 70 to 100 mg per cent and are presumably very close to the true glucose values. The specificity for glucose appears to reside in the method of deproteinization, since filtrates prepared as described may be analyzed for glucose by the Folin-Wu or Benedict colorimetric methods to give results substantially identical with those obtained by the Shaffer-Hartmann rodometric titration. For the principle of the Shaffer-Hartmann procedure, see Chapter 31

4. Nelson-Somogya Method ** Principle. Blood is deproteinized by a zine hidroxide-barium sulfate procedure which gives a fittate containing practically no reducing substances other than glucose. The zine-barium filtrate is heated with an alkaline copper reagent and then treated with a special arisenomoly bladice color reagent. The color developed is compared with that obtained from a known amount of glucose.

Procedure. (a) Derisories ization Place 1 ml. of blood in a 50-ml. flask. Add 9.5 ml. of barium hydroxide solution, mixing by rotation. Add 9.5 ml. of

[&]quot;Nelson J Biol Chem 153, 375 (1944), Somogo: J Biol Chem., 150, 62 (1945). Adapted by permission from the official procedures of the American Association of Clinical Chemists.

^{**} Reagenta Required Barsum Hydroxide Solution Dissolve 90 g of Ba(OH); Slf;0 in distilled water and dilute to 2000 ml in a graduated cylinder I ilter if cloudy Store in well-

574 zinc sulfate solution, mixing by rotation Shake vigorously and filter on a dry filter paper, collecting the filtrate in a dry fiask

(b) DETERMINATION OF GLICOSE Measure 0 5 ml of the barlum-zine filtrate into a test tube calibrated at 10 ml Add 1 ml of alkaline copper reagent, mix -hy tapping cover the top of the tube with a marble, and place upright in a boiling water hath for 20 minutes Cool by placing the tube in water at room temperature for I minute Add I ml of arsenomolybdate color reagent, mix ing by tapping then dilute to 10 ml with distilled water. Mix by inversion Compare in a colorimeter against a simultaneously prepared standard oh tained by treating 0.5 ml of standard glucose solution (see footnote 90) exactly as described for the blood filtrate For photometric measurement prepare a blank by carrying out the procedure as described on 0.5 ml of water Set the photometer to zero density with the blank, and measure the optical densities of standard and unknown at 540 mu

CALCULATION For colorimetrie n eas tren ent calculate as follows

Reading of Standard \times mg glucose in stan lard $\times \frac{100}{0.025}$

- mg glucose per 100 ml blood

It is convenient to set up three slandards covering low normal and high values of blood glucose as follows 1 containing 0 0120 mg glucose for low values 11 con taining 0 025 mg glucose for normal values 111, containing 0 050 mg glucose for high values The unknown is then compared against the standard it most closely

stoppered containers filled to capacity

Zine Sulfale Solution Dissolve 100 g of InSO, 711:O in distilled water dilute to 2000 ml

in a graduated cylinder and mix. Statte indefinitely

It is most important that these two solutions exactly neutralize each offer Check b) measuring 10 ml of zinc sulfate solution into a flask diluting with about 50 inl of distilled water add 4 drops of prenolphthalein solution and titrate with the harium) ydrox de solution slowly and with constant shaking until t drop of alkali turns the maxture p nk If it requires more or less than 10 00 ml ± 0 05 ml of barium hy droxide dilute one or the other solution as required and cleck again Continue until the two solutions are exactly equivalent Protect il e barium hydroxide solution in the dispensing bottle with a soda-lime tube in the stopper and test both solutions occasi nally to making a trial filtrate on blood Satisfactory solutions will give a clear filtrate which filters rapidly and shows little tend enes to foam

Alkaline Copper Reagent Solution A Dissolve 50 g of anhydrous acdium carbonate 50 g of Roci cle sait 40 g of sodium bicarbonate and 400 g of anhydrous sodium sulfate in about 1000 ml of distilled water and d tute to 2 litera Mix and filter if not clear Store at room temperature If a sediment forms in a few days filter again Solution B Dissolve 150 g of CuSO, 5thO in dist lied water and d tute to I liter Add 0.5 ml of concentrated sul in current outs in ustrice water and dute to 1 liter Add 0.5 ml of concentrated surfure and and mix. Alkaline Copper Respect. On the day it is to be used place 4 ml of colution B ma 100-ml graduated cylinder 4 lite to 100 ml with Solution A and mix. Attenomolybelate Color Peoper Divisions 100 of committee of aminonium molybelate in 1800 ml of datafilled water Add 84 ml of concentrated authorities.

disodium orthogramate (\ail\last1\last1\last2\tau\TH_0) in 100 ml of distilled water and add it with attring to the acid fied molybdate solution. Place the mixture in an incubator at 3 of for I to 2 days Store in a glass-etoppered brown glass bottle. This solution should be stable

Standard Glucose Solutions Prepare a stock standard by dissolving exactly 100 g of lighest purity and drous glucose in about 10 to 15 ml of 0.2 per cent benzo c acid sold tion transferring quantitatively to a 100-ml volumetric flask diluting to the mark with the benzon acid solution and mix ng This solut n is stalle indefinitely and contains 10 mg of glucose per ml Standard 1 (0 0125 mg, glucose per 0 5 ml) is prepared by diluting 0.5 ml of the stock at and ard to 200 ml with 0 2 per cent benzo e and solution and mixing For Standard II (0 020 mg glucose per 0 5 ml) dinter 10 ml of stock standard to 2012 20 ml of stoc mi of stock standard to 200 ml Tiese d lute solutions in 0 2 per cent benzoic acid solution keep indefinitely at room temperature

matches. The calculation can be simplified as follows

 $\frac{\text{Reading of Standard}}{\text{Reading of Unknown}} \times f = \text{mg glucose per 100 ml blood}$

where f is a factor with the following values for standard I, 50, for standard II, 100, for standard III, 200

For photometric measurement the calculation is as follows

 $\frac{\text{Density of Unknown}}{\text{Density of Standard}} \times \text{mg glucose in standard} \times \frac{100}{0.025}$

= mg glucose per 100 ml blood

A standard containing 0 025 mg of glucose is satisfactor; for blood samples containing up to 350 mg per cent blood sugar With this standard the calculation simplifies to

Density of Standard × 100 = mg glucose per 100 ml blood

For samples containing more than 350 mg per cent blood sugar, repeat the analysis using a smaller aliquot of filtrate plus water to make 0.5 ml and correct the calculations accordingly. The colors obtained by this method are said to be stable for more than 24 hours.

Interpretation. Normal values on fasting venous blood by this method range between 65 and 110 mg per 100 ml of whole blood, with an average value of 85 mg per 100 ml. The interpretation of variation from the normal range is similar to that already presented in connection with the Folin-Wu method.

5. Micromethod for Glucose in 0.1 Mi of Blood (Folin and Malmros) ²¹ Principle The sugar is oxidized with alkaline potassium ferricy anide and the ferrogranide produced is measured colorimetrically or photometrically after conversion to Privasian blue

Procedure With an accurate 0 1-mi pipet" collect 0 1 ml of blood and transfer it to 10 ml of dilute tunestic acid" in a centrifude tube. Str. well and

[&]quot;I Folia and Malmros J Biol Chem 83 115 (1929) Folia J Biol Chem 77, 421 (1928) See also Horvath and Knehr J Biol Chem 148 869 (1941) For a method based upon uttration of the ferrory and formed using cerns sulfate see Miller and Van Slyke J Biol Chem 14 583 (1936) MacFadyen and Van Slyke third 149, 527 (1943) See also the Hawkins and Van Slyke method Chapter 31

^{*1} A capillary pipet about 7 cm in length per 0 1 ml calibrated to contain using mer-

cury (0 i ml. weighs 1.355 g.) Sold by dealers in laborators supplies

**Pollute Tungstic Acid Solution Transfer 20 ml of 10 per cent sodium tungstate to a liter
volumetric flask. Dilute to about 800 ml. Add with shaking 20 ml. of 25 N sulfuric acid and

dilute to volume

Potassum Ferregande Solution Dissolve 2 g of c p potassuum ferregande in distilled
water and dilute to a volume of 500 ml Keep the major part of the solution in a brown bottle in a dark closet Keep the rezizent in dails, use also in a brown bottle

Sodium Cyanude-Carbonate Solution Transfer 8 g of anhy drous sodium carbonate to a 500 ml volumetrie flask Add 40 to 50 ml of water and shake to promote rapid solution With a cylinder add 150 ml of freshly prepared 1 per cent sodium cyanide solution didute to volume and mix

Ferrie Iron Solution. Fill a liter cylander with water. Suspend on a copper wire screen just below the surface 20 g of soluble guing shaits and leave overnight (18 lours). Remove the screen and strain the liquid through a double layer of a clean towel. Add to this extract a solution of 5 g of anh glorous ferrie sulfate in 75 m lot 85 per cent phospl ore acid plus 100 ml of water (1dd to the mixture a little at a time about 15 ml of 1 per cent potassium permanginant solution to destroy certain reducing materials present in ginn glatti. The

576 centrifuge. Transfer 4 mt of the water-clear supernatant fluid to a test tube graduated (with a ring going all around) at 25 ml. Transfer 4 ml. of the standard sugar solution to another similar tuhe. To each tulie add 2 ml, of the 0.4 per cent potassium ferricyanide solution and 1 ml. of the cyanide-carbonate solution. Heat immediately in holling water for 8 minutes, and cool In running water for 1 to 2 minutes. Add 5 ml. of the ferric iron solution and mix. Let stand for 1 to 2 minutes and then dilute with water nearly, but not quite, to the 25-ml, mark. Add 2 drops of alcohol to cut the foam and dilute to the mark, Mix. Allow to stand 10 minutes and read within the next 30

For colorlmetric comparison, half fill the colorlmeter cups with the greenminutes. colored standard, set the two plungers at a height of 20 mm., and cover the opening of the light box with the pictic acid light filter. 4 Adjust the position of the colorimeter and of the mirror glass reflector until the two fields look exactly alike. The adjustment is easier if the colorimeter is kept on a piece of plate glass polished on one side and rough on the other. Rinse one colorimeter cup and plunger with the unknown solution and pour the unknown into the cup to a sultable height. Compare in the usual way.

For photometric measurement, transfer the solutions to suitable containers and determine the densities in a photometer at 520 mg. Set the photometer to zero density with water alone, rather than use a hlank (see helow).

CALL UI ATION For colorimetric measurement

Reading of Standard Reading of Unknown
$$\times 0.04 \times \frac{10}{40} \times \frac{100}{0.1} = \text{mg}$$
 glucose per 100 ml blood

The proportionshity between color and concentration is excellent in this method so that readings between 5 and 40 mm may be accepted, with the standard at 20 mm, if perfect equality as to light and color is obtained If the blood glucose concentration is over 100 mg per cent, repeat with 2 ml of filtrate plus 2 ml of water, and multiply the results by 2

For photometric measurement

Density of Standard
$$\times$$
 0.04 \times $\frac{10}{40}$ \times $\frac{100}{0.1}$ = mg glucose per 100 ml blood

At the wavelength specified and in a 1-cm cuvette, the standard corresponding to

slight turbidity of the solution will disappear completely if kept at 37° C for a few days

The use of Duponol a synthetic detergent has been recommended in place of gum ghatti (Klendshoj and Hubbard J Lab (In Wed 28, 1102 (1939 1940) Horvath and Knehr

Standard Glucose Solution The working standard contains 0 01 mg of glucose per ml (see footnote 78 p 568)

"Heating for 20 minutes instead of 8 minutes to ensure complete exidation of glucose

has been recommended by Jourdonaus J Lab Clin Med 23, 847 (1937-1938), and by Horvath and Knehr loc cu ** Dissolve 5 g of pierie acid in 100 ml of methyl alcohol and add 5 ml of 10 per cent

NaOII Place a pack of 8 to 10 heavy absorbent filter papers (Schleicher and Schüll No is good) on a level and smooth mat of newspapers. Pour the acid picrate solution over the filters until the papers are saturated and an excess of solution filters through Allow to dry When perfectly dry pour over the pack an excess of solution filters through Allow to order when perfectly dry pour over the pack an excess of a 3 per cent solution of parafin of bennine. Allow to dry All papers should be evenly stained. Cover the light box window of the colorimeter lawn with stands of sections. the colormeter lamp with staned paper. Test the efficiency by comparing water and at the centre of the colormeter lamp with staned paper. Test the efficiency by comparing water and at the centre of two fields look alike II the filter is madequate equality of the fields cannot be obtained A

The MEMORIAN SINK II the Bifer is undequate equality of the fields cannot be obtained from kight is necessary. A filter in the form of a glaw duk to be put on top of the occular say be obtained from the Klett Manufacturing Co. New York See Tauber J. Leb. Clin fed. 18, 760 (1930)

fed 15, 766 (1930)

100 mg per cent blood glucose has a density of about 0 250 permitting accurate measurement up to about 400 mg per cent blood glucose For higher values or with deeper cuvettes use 2 ml of filtrate plus 2 ml of water in the analysis and multiply the results by 2. With well prepared reagents, the blank should have a density of less than one-tenth that of the standard given in the presence of a blood filtrate the blank density is even less and better agreement with Beer's law is noted if the blank is neglected and the photometer set to zero density with water alone as described 95

Interpretation. Normal values by this method range from 75 to 105 mg per cent blood glucose It must be noted that, if fingertip blood is used, this type of blood is essentially arterial, and resembles venous blood only in the fasting state. Thus during absorption or in a glucose tolerance test higher glucose values will be obtained on fingertip blood than for venous blood, although the reverse may obtain in severe diabetes For further interpretation, see p 570

6 Determination of Sugar (Method of Hagedorn and Jensen) 97 Principle The blood protein is precipitated with zinc hydroxide. The filtrate is heated with potassium ferricy anide solution and the amount of ferricy anide reduced is determined by adding an iodide solution and titrating the iodine set free with sodium thiosulfate The principal reaction is 2H₂Fe(CA)₆ + 2HI = 2H₄Fe(CN)₆ + I₂ The reversal of the reduction reaction is prevented by precipitation of the ferroeyanide formed as a zinc salt

Procedure " Into a test tube (15 × 150 mm) pipet 1 ml of 0 1 N NaOH and 5 ml 0 45 per cent zlac sulfate solution A gelatinous precipitate of zlac hydroxide forms 0 1 ml of blood from a capillary pipet " is introduced, the

⁹⁶ Unpublished results of Summerson and Robinson

⁴⁷ Hagedorn and Jensen Brochem Z 135 46 (1923) and 137 92 (1923) For critical stud ies of this method with suggested improvements see Polin and Malmros J Biol Chem 83

121 (10°0) Kramer and Steiner Brochem J 25 161 (1931)

** Reagents Required For Protein Precipitation 01 h haOH - Zinc sulfate 0 45 per cent These colutions are best prepared every eight days by dilution of 2 N NaOH and a zinc sul fate solution containing 40 g of the salt in 100 ml of solution For Sugar Determination Potassium ferricyanide 1 65 g and sodium carbonate (fused) 10 6 g in 1000 ml of water Protect from light Iodide-sulfate-chloride solutions KI 5 g zinc sulfate 10 g NaCl 50 g water to make 200 ml It is best to prepare the solution without iodide and add the latter to portions of the solution as required. Free jodine can be almost completely removed by filtering through thick paper. The blank will take care of smaller errors. Acetic acid solution. 3 ml of acetic acid (iron free) with water to make 100 ml Starch solution 1 g of soluble starch dissolved in 100 ml of saturated NaCl solution Sodium thiosulfate solution 0 " g sodium thiosulfate in 500 ml of water 0 000 A Polassium todate This solution is permanent and is used to cleck the thiosulfate and ferries ande solutions which keep less well. It slone need therefore be prepared very accurately. Dissolve 0.3506 g. potassium iodate (water free) in water to make 2009 ml

Chemicals of lighest purity must be used Sodium carbonate is best recrystallized and fused in platinum. Acetic acid should be tested for iron, as should also the zinc sulfate sodium chloride and potassium rodide. The icdide should also be tested for rodate. Ordinary quantities of the mixed solutions will then give no test for iodine with starch but will give a test if 0.01 ml of ferrica anide is added. Potassium ferricy anide is prepared by washing ery stals of the ordinary product with water dissolving in water with heat and filtering the boiling solution through paper previously carefilly washed with boiling water into an evaporating dish set in ice-cold water Tle fine crystals are filtered off with suction on another washed paper and again reery stallized Dry at 50° C. Keep away from sunlight during course of preparation

" The length of the 0.1 ml pipet from the tip to the mark should be about 10 to 12 cm and over-all length about .0 cm Calibrate as follows With the pipet measure out 0.1 ml of 0.1 N potas i un iodate solution into 10 i il of water. Add acid and KI solution in the usual way and titrate with 0.02 N (I osulfate. Some of the same iodate solution is then dilute I to 0.02 \ and exactly 2.0 ml of this in 10 ml of water titrated with the same thiopipet being washed out twice with the mixture and blown empty Put in a boiling water bath for 3 minutes Filter on a funnel of 3 to 4 cm. diameter, prepared with a small filter of washed, moistened, not tightly pressed cotton, into a test tube (30 × 90 mm) Wash the funnel and filter with two 3-ml portions of water Add 2 ml of alkaline potassium ferricyanide solution and heat in a hoiling water hath for 15 minutes Cool and add 3 ml of the lodide-sulfate solution and 2 ml of 3 per cent acetic acid solution. Titrate with 0 005 N sodium thiosulfate, using as an indicator 2 drops of 1 per cent solution of soluble starch in saturated sodium chloride solution

CALCULATION Determine the blank obtained by carrying through the whole determination but without the addition of blood 100 Multiply A (the thiosulfate burst reading) by the factor for the thiosulfate (2 00/ml thiosulfate required for 2 ml 0 005 N todate) Express this value (B) for unknown and for blank as mg glucose by consulting the table which follows Subtract the glucose value of the blank from the glucose value of the unknown. The difference is mg. glucose in 0.1 ml. of blood. Or cal culate as follows 2 00 - B = z (ml 0 005 N ferries anide reduced) 2 00 ml of ferries cyanide are reduced by 0385 mg of glucose Therefore mg per cent glucose = 0385 $\times \frac{x}{2.0} \times 1000$

Interpretation. Similar to previous method

MILLILITERS 0 005 NatSiO: Used and MILLIGRAMS GLICOSE PRESENT

MI
00 01 02 03 04 05 06 07 08 09 11 11 12 13 14 15 16 17 18

7. Manometric Methods for Reducing Sugars in See original papers

sulfate as before The two titrations should agree within experimental error (about 0.5 per ent) if the pipet is accurate

¹⁴⁴ Actions and \$1 yearsybutyris and do not reduce the reagent. One mg of uric acid aves a reduction equal to that 10 53 mg of glucose and 1 mg of creatinine a reduction equal to 0 47 mg of creatinine a reduction

¹⁰¹ Van Elyke and Hawkins J Biol Chem 79 733 (1928), 83, 51 (1929)

CARBOHYDRATE TOLERANCE TEST

Principle. Blood sugar is determined at hourly periods following the ingestion of 1 g. of glucose per kilogram of body weight. Urinary sugar for the 24-hour period following the ingestion of the glucose is also determined.

Procedure. The first thing in the morning, collect a specimen of urine and one of blood to serve as controls. Then give the patient 1 g. of glucose per kilogram of body weight. The glucose may be given in 50 per cent solution, Collect three or four specimens of blood at hourly intervals and analyze for sugar. Following the taking of glucose collect a 24-hour specimen of urine and determine its sugar content. Brillier and others have proposed the use of a test breakfast instead of the glucose meal.

Interpretation. In normal individuals, blood sugar rises from the normal value of about 100 mg, to about 150 mg per 100 ml, at the end of the first hour, and returns to normal by the end of the second hourly period. In pathological conditions, the curve does not follow the normal course. Hyperthyroidism, diabetes mellitus, and nephritis show much greater values, depending on the severity of the disease, and the return to normal is delayed for three hours or more. The higher sugar concentration in the blood during the test may or may not he accompanied by glycuresis, depending upon the "threshold point" of the kidney. In diabetes the threshold point is usually, and sometimes markedly, increased above its normal range of 160 to 180 mg per 100 ml. In hypoendocrine conditions, in which the blood sugar is low ordinarily, the curve of blood sugar during a tolerance test is quite flat. Wherever possible, it is hetter practice to ohtain quarter-hour specimens of blood, during the first hour. This provides more information concerning the general nature of the curve and the height of the peak. John 103 helieves that a diagnosis of diabetes is justified if the curve remains high after three hours, irrespective of the height to which it has risen. Petty and Stoner104 on the other hand regard the criterion as a rise above 180 mg. They have their claim on simultaneous determinations of respiratory quotients, which show very slight rise in diabetics, but follow the sugar curves in normal individuals and in renal glycosuries. In nondiabetics the peak of the curve is usually between one-half and one hour, while in diahetics it is nearer two hours.

It has been shown by Foster 105 that earbohydrate tolerance tests conducted on finger blood, which is practically arterial, give higher and sharper curves than tests of venous blood collected simultaneously. It appears that some of the glucose of arterial blood is removed and oxidized or stored as glycogen by the tissues The various mechanisms for glucose utilization appear to be stimulated to overactivity, since the blood sugar

¹⁰¹ Brill J. Lab Clin. Med , 8, 727 (1923).

¹⁴² John Am J. Med. Sci., 169, 102 (1925). J. Metabolic Research, 1, 497 (1922).

Petty and Stoner: Am. J. Med. Sci., 171, S42 (1925).
 Hamman and Hurschman. Arch. Internal Med., 20, 751 (1917); Bailey. Arch. Internal. Med., 23, 455 (1919); Williams and Humphreys: Arch. Internal. Med., 23, 557, 546, 559 374. A. A. A. G. (1919); Whinains and Humphreys. Rect. Activation for the Medical Research, No. 11, 1919, Macleod: Physiol. Rets. 1, 208 (1921); Foster. J. Bol Chem. 55, 291, 303 (1923); Du Vigneaud and Karr. J. Biol. Chem. 64, 281 (1925); Hubbard and Wright; Clyson Med Bull., 12, 155 (1926); Rabinowitch: Bril. J. Expl. Path., 8, 76 (1927).

curve frequently falls below its initial level. For further discussion of the application of sugar tolerance tests consult papers by Hamman and Hirschman, Hubbard and Wright, Badey, Williams and Humphreys Allen, Stillman and Fitz, Rabinowitch, Macleod, Foster, and du Vigneaud and Karr (See references in footnote 105)

DETERMINATION OF CHOLESTEROL

Introduction. The cholesterol of blood is present in the form of both free cholesterol and cholesterol esters. In the plasma both free and esterified cholesterol is found, in the red cells only the free form appears to be present. Plasma is preferred to whole blood for analysis, since pathological variations in the amount and in the distribution between free and ester forms occur largely in the plasma fraction. Free cholesterol is best determined by precipitation and isolation as the insoluble digitation, or otherwise determined on the trace of the precipitation and isolation as the insoluble digitation, the precipitate may be weighted, or treated with a cholesterol color reaction, or otherwise determined. Total cholesterol is most accurately determined in a colorimetric procedure after preliminary saponification to liberate ester cholesterol from combination, since free cholesterol and ester cholesterol may give different color intensities per unit amount of cholesterol present. Many methods which do not involve saponification bave, however, been proposed, as discussed below.

1. Method of Schoenheimer and Sperry ** Principle. An acctone-alcohol mixture is used to precipitate the proteins and extract the cholesterol and cholesterol seters from the sample of whole blood or (preferably) plasma or serum. The cholesterol is precipitated with digitonin, either before asponification (free cholesterol) or after asponification (total cholesterol) and the separated digitonide is purified and subjected to the Liebermann Burchard color reaction. The color is then compared with that produced by a standard cholesterol solution.

Procedure 1st Place approximately 10 ml of the acetone-alcohol mixture in a 25-ml glass stoppered volumetric flask and add 1 ml of serum or plasma down the side of the flask below the graduation After withdrawing the piper,

¹⁸⁸ Schoenheimer and Sperry J Biol Chem 106 745 (1934) Sperry Am J Clin Pelh 8 Tech Suppl 2 91 (1938) Sperry (personal communication) See also Fitz J Biol Chem 107 523 (1935) Sobel and Mayer J Biol Chem 137, 255 (1945) The original Schoenheimer-Sperry method is a microprocedure requiring but 02 ml of sample Theorecute described here is a shalthy modified version which uses a larger sample require less technical skill and has been found satisfactory for routine use in several laboratories. If Research I Regents Required Actions Alcohol Matter's Unit 2 volume of absolute cityl slicing.

with 1 volume of redistilled accione
Distribution Solution Dissolve 400 mg of digitomin (Hoffman LaRoche Nutley N J or
S B Pemek and Co 50 Church St. New York N Y) in 100 ml of distilled water Fifter

or centrifuge just before use if not clear

10 Per Cent Actic Acid Solution Dilute 1 volume of glacial acetic acid to 10 volumes with
water and mix

Accione Ether Mixture To 1 volume of redistilled accions add 2 volumes of peroride-free ether and mix To prepare peroxide-free ether wash ordinary ether with sodium suffice followed by water and distill from calculum chloride

Potassium Hydroxide Schidon Davide 10 g of reagent-grade potassium hydroxide in 20 m of water Store in a boottle reupped with a med cine dropper earry ing a rubber blue Phenolphthalein Schidon Davide I g of phenolphthalein in 65 per cent alcohol and dute to 100 ml

Acric Acid Only the highest purity anhydrous product (giscial) may be used Acric Anhydride Sulfuric Acid Reagent Just before needed place 20 ml of acetic anhy

swirl the flask to produce a finely divided precipitate, then immerse the flask in boiling water, with swirling to prevent bumping, until the solvent boils. Remove, cool to room temperature, and make up to volume with the alcoholacetone mixture. Stopper, mix thoroughly, and pour onto a dry filter, collecting the clear filtrate in a dry test tube. Cover the funnel with a watch glass during filtration, to minimize evaporation of solvent.

Precipitation of Free Cholesterol, Transfer 6 ml. of filtrate to a 15-ml. centrifuge tube, add 3 ml. of digitonin solution, and 1 drop of 10 per cent acetic acld solution. Place a stirring rod in the tube and stir thoroughly. Place the tube, together with the stirring rod, in a pint or quart size preserving jar, cover the jar tightly, and allow to stand at room temperature overnight, in the morning, transfer the tube and rod to a test tube rack and stir cently to free particles of precipitate which may have adhered to the walis of the tube, Remove the rod carefully without touching the upper part of the tube and place it aside carefully so that no adherent precipitate is rubbed off; a wire rack with numbered positions is suggested, so that the rod may be returned to the proper tube later. Centrifuge the tube for about 15 minutes at about 2800 r.p.m. Decant the supernatant and drain in an Inverted position for a few minutes, removing the last drop by touching the lip of the tube to a clean towel; or remove the supernatant by geotle suction through a fine capillary pipet, without stirring up the precipitate or touching the sides of the tube. A few particles may float at or near the top of the solution after centrifugation; they are discarded with the supernatant.

Replace the stirring rod in the tube and wash down the walls of the tube and the rod with 1.5 to 2 ml. of acetone-ether mixture added from a dropping pipet with attached rubber bulb. Stir the precipitate up thoroughly, replace the rod on the rack, and centrifuge the tube for 5 minutes. Remove the supernatant as above, and repeat the washing twice more, using ether instead of acetone-ether. Replace the rod in the tube and set aside until ready for color development.

Saponification and Precipitation of Total Cholesterol. Add 3 drops of potassium hydroxide solution to a 15-ml, graduated centrifuge tube, and add 3 ml. of the acetone-alcohol filtrate. Insert a stirriog rod and stir vigorously until no droplets of aikall can be seen in the solution. Leave the rod in the tube, and place tube and rod in a preserving jar containing a layer of sand about 3 cm, deep which has been previously heated in a water bath until the temperature of the sand is about 45° C. The sand acts as a heat reservoir. Glose the jar tightly and place in an incubator (37° to 40° C.) for 30 minutes Remove the tube to a rack, allow it to cool to room temperature, raise the rod, and add alcohol-acetone mixture to the 6-ml. mark. Add 1 drop of phenolphthalelo solution, followed by 10 per cent acetic acid, drop by drop

drude (Eastman) in a glass stoppered eximder and chill in rec water. When cold add 1 ml of concentrated sulfure acid, a little at a time, with mixing and cooling during the addition. Stopper the cylinder, shake the contents vigorously for a few moments and return to the ice both. Keep cold during use, and do not use any reagent more than 1 hour old. More or less of the respect may be prepared as needed, using the same proportions.

Stock Cholesterol Standard Dissolve 100 mg of pure dry cholesterol in about 50 ml of glanda actic nead by warming on an electric hot plate and stirring Transfer with rinsings to a 100-ml glass stoppered volumetric flask dilute to 100 ml with acetic acid, and mix This solution contains I mg of cholesterol per ml and is quite stable in the cold it in uset be warmed to room temperature before using Dilute standard Transfer 5 ml of stock standard containing 5 mg of cholesterol, to a 50-ml glass-toppered volumetric flask dilute to mark with sectic acid, and mix well. This solution contains 0 2 mg of cholesterol is

with stirring, until the red color disappears Add 1 drop in excess, followed by 3 ml. of digitonin solution. Stir thoroughly, place the tube and rod in a preserving jar, cover tightly, and allow to stand at room temperature for at least 3 hours, and preferably overnight. Separate and wash the precipitate exactly as described above for the precipitation of free cholesterol, except that only one ether washing instead of two is necessary.

Development ond Reoding of Color. Place a layer of sand about 3 cm. deep in a shallow pan and heat to 110° to 115° C. In an oven. Place the tubes containing the precipitated and washed cholesterol digitoride in the pan and return the pan to the oven. ** After 30 minutes, remove the pan and, while the tubes are still in the hot sand, from a buret add 2 ml. of acetic acid in such a way that the acid washes down the wail of the tube and the rod. Stir well with the rod, allow to remain in the sand for about 2 minutes (not longer), remove and cool to room temperature, with the rod still in the tube. Undissolved material at this stage is neglected.

When the tubes are ready for color development, place them in a water bath at 25°C from which light is excluded 100 and allow to come to temperature equilibrium. Note the time, and add 4 ml of the cold acetic anhydride sulfuric acid reagent, mix well with the rod, and return to the bath Allow to atand 27 minutes, then transfer to colorimeter or photometer containers and read within the next 10 minutes. If possible, it is better to have the time of standing before reading constant at about 30-31 minutes

A standard suitable for both colorimetric and photometric measurement is prepared as follows. Transfer 2 ml of a standard solution of cholesterol in acetic acid, containing 0.2 mg of cholesterol, to a suitable tube containing a stirring rod. Place in the water bath at 25° C, and allow to come to temperature equilibrium. When ready, add 4 ml of the acetic anhydride-suifuric acid reagent, mix, return to the bath, and allow to stand 27 munites. Use within the next 10 minutes. We rephotometric measurement, a blank is prepared similarly, except that 2 ml. of acetic acid alone is used instead of the standard cholesterol solution.

For colorimetric measurement, the colorimeter must be equipped with microcups and the eyeplece of the colorimeter must be provided with a red filter (Wratten No. 71A, supplied by Eastman Kodak Co or by the manufacturer of the colorimeter). Arrange the time of color development so that the standard is ready first, and the unknowns reach the 27-mmute stage at short intervals thereafter, depending upon the rapidity with which the color comparisons can be made. Adjust the standard against Itself at 20 mm, and as each unknown becomes ready, rinse cup and plunger with a small portion and read against the standard Do not use the standard after 37 minutes have elapsed from the start of color development; if more unknowns are present than can be read in this time, have a second standard suitably timed and ready for use ""

¹⁰² This heating is to ensure complete removal of water from the hygroscopic digitonide precipitate.

¹⁹ A large pan of water containing metal racks and placed in an enclosed box fitted with a door or in a closet is satisfactory Insert a thermometer in the water and keep at 25° C by adding hot or cold water as required.

¹⁰ Because the standard appears to fade rapidly while it is in the colorimeter cup Fits (or crt) suggests the preparation of a larger volume of standard fresh portions of which are used at intervals during the 21 to 37 minute period of color development.

because of the install day of the standard spend of cours of evelopments on a number of analyses, are being run colormentrically artificial been suggested that when a number of analyses, are being run colormentrically artificial standard level each day against a cholesterol standard Shapiro Lerner and Powen (Free See Erst Biol Vied. 23, 1300 (1933)) proposed a permanent artificial standard wade from Genter and Course of the Cours

For photometric measurement, transfer the colored solutions to suitable containers and determine the density in the photometer, at 625 m μ Adjust the photometer to zero density against water, and determine the densities of the blank, the standard, and the unknowns at 30 to 31 minutes after adding the acetic anhydride sulfuric acid reagent. Subtract the density of the blank from those of the standard and unknowns, to obtain the true densities.

CALCULATION For colorimetric measurement. Since the amounts of filtrate taken for the free and total cholesterol determinations represent 0.24 and 0.12 ml of original sample respectively, the calculations are as follows.

Free Cholesteral

Reading of Standard Reading of Unknown
$$\times 0.2 \times \frac{100}{0.24}$$
 = mg free cholesterol per 100 ml

Total Chalesteral

$$\frac{\text{Reading of Standard}}{\text{Reading of Unknown}} \times 0.2 \times \frac{100}{0.12} = \text{mg total cholesterol per 100 ml}$$

For plotometric measurement

Free Chalesteral

$$\frac{\mathrm{Density} \ \mathrm{of} \ \mathrm{Unknown}}{\mathrm{Density} \ \mathrm{of} \ \mathrm{Standard}} \times 0 \ 2 \times \frac{100}{0 \ 24} = \mathrm{mg} \ \mathrm{free} \ \mathrm{cholesterol} \ \mathrm{per} \ 100 \ \mathrm{ml}$$

Total Cholesterol

Density of Unknown
$$\times 0.2 \times \frac{100}{0.12} = \text{mg}$$
 total cholesterol per 100 ml

At 025 mµ and in a 1-cm cuvette the density of the standard is approximately 0 100 m (Fig. 153). Under the conditions described from 40 to 200 mg per cent free cholesterol and 80 to 400 mg per cent total cholesterol may be accurately determined.

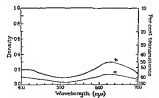


FIG 153 ABSORPTION SPECTRA OF COLORED SOLUTIONS OBTAINED BY SCHOCKHEIMER SPERRY METHOD FOR CHOLESTEROL.

FOR standards containing (a) 0.2 mg and (b) 0.4 mg cholesterol Solution denth 1 cm

Midinght Black Ink. Sperry and Brand (J. Biol Chem. 150, 251 (1943)) describe in detail the use of dilute napl it of green B solution as an artificial standard.

¹¹¹ Color developed for 30 to 33 nunutes at 21° C

For higher levels, repeat the analysis using less sample in the alcohol acctone precipi tation or use less filtrate in the digitonin precipitation with other reagents in proportion except for the final solution in acctic acid and color development, and correct the calculations accordingly

Because of the sensitivity of the clolesterol color to bleaching by light photometric measurements should be made as rapidly as possible to avoid prolonged exposure to the light beam in the photometer

Interpretation. Normal blood cholesterol appears to be maintained at a constitutional level which is characteristic for each individual and from which large deviations do not ordinarily occur for that particular individual Considerable variation however, is found among different individuals the normal range being from about 110 to 390 mg per 100 ml of serum or plasma Of this, about one-third is present as free cholesterol (on the basis of digitonin precipitation, which is generally accepted as standard) and the remainder is esterified. Although many of the data in the literature are in terms of total cholesterol, the distribution of cholesterol between the free and the ester forms is attracting more attention, particularly in the diagnosis of liver disease. According to Sperry, routine determination of whole blood cholesterol should be aban doned, since the cells contain only free cholesterol and in quite constant amount no changes in pathological conditions having been noted. Scrum cholesterol is increased and the determination has clinical value in nephrosis lipemia diabetes mellitus, hypothyroidism and biliary obstruction due to calculi or other causes Increases are also found in preg nancy and after a high lipide diet Decreased serum cholesterol is found in hyperthy roidism, pernicious anemia, and certain types of liver disease In the latter condition, the proportion of ester cholesterol to total cholesterol may be sufficiently lowered to be of diagnostic significance. A low choles terol diet leads to lowered serum cholesterol, since there is ample evidence that cholesterol is synthesized within the hody changes in blood level due to diet presumably represent merely the net result of the various factors concerned

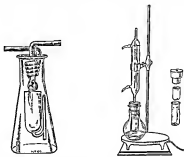
2 Resphold and Shiels' Modification of the Myers-Wardell Method Principle The serum or plasma is dried on anhydrous sodium sulfate and extracted with chloroform The total cholesterol of the extract is determined colorimetrically by the Liebermann Burchard reaction with acetic ambydride and sulfuric acid. In the original method of Myers and Wardell plaster of Paris was used instead of sodium sulfate Leiboff¹¹⁴ describes a procedure in which the sample is dried on a small piece of absorbent paper which is then extracted in a specially designed flask. In the Bloor Pelkan and Allenns procedure no special apparatus is necessary proteins are precipi tated with alcohol-ether mixture an aliquot of the filtrate is dried in a beaker and ex tracted with successive small portions of hot chloroform. In all of these procedures low results will be obtained if the extraction is incomplete

Bloor leikan and Allen J Biol Chem 52 191 (1922) See also Kelsey J Biol Chem 127 15 (1939)

¹¹¹ Myers and Wardell J Biol Chem 36 147 (1918) Reinhold and Shiels Am J Clin

Path 6 27 (1931) 114 Leiboff J Biol Chem 61 177 (1924) J Lad Clin Wed 10 8.7 (1925) 11 77 (1926) 15 776 (1930)

Procedure 114 Transfer I ml of plasma or serum to a small mortar contain ling about 8 g of anhydrous sodium sulfate Mix uniformly, dry in an oven at 100°C for 10 minutes, cool in a desiccator, pulverize, and transfer completely to a paper extraction shell which is then inserted into a Soxhlet extraction tube (see Fig. 154) suspended from the condensing coll of the extraction apparatus (if the extraction apparatus illustrated in Fig. 154 is not avail able, Myers' arrangement shown in Fig. 155 may be used, in which the extraction tube is a perforated glass tube (25 × 7 cm) and is connected to



FIGS 154 AND 155 EXTRACTION APPARATUS FOR CHOLES-TEROL DETERMINATION

Fg 100 after Myers P actical Chemical Analysis of Blood 2d ed C V Mosby Co St Louis

a reflux condenser as illustrated) Place 20 to 25 ml of redistilled chloroform in the extraction flask, place on an electric hot plate, and extract for 90 mln tutes, during which time cold water flows through the condenser or coil Allow the extract to cool and transfer with rinsings to a 25 ml volumetric flask. Make up to volume with chloroform, mix well, and filter through a dry filter if necessary.

Transfer 10 ml of the chloroform extract to a dry test tube and add 2 ml of the freship prepared acetic anhydride-suffuric acid reagent Treat 10 ml of a standard solution of cholesterol in chloroform, containing 0 8 mg of

¹¹⁶ Reagents Required Anhydrous sodsum sulfate reagent grade

Chloroform. Commercial chloroform abould be weahed with water dried over anhitdrous potassium carbonate and distilled Dr. He distillate over phosphorous pentoxide and ogain distil. Acep protected from light. Present-day reasent grade chloroform (such as Merck s) may be satisfactory with out it e necessity of furtler treatment.

may be satisfactory with out the necessity of further treatment.

Acetic Anhydri le-Sulf ric Acid Reagent Just before use add 1 volume of concentrated sulfuric acid slowly with all slung to 10 volumes of acetic and dride. Prepare only enough

reagent for a particular series of analyses and d-seard it e unused portion. Cholesterol 5th adra? Stock Standar? Dissolve 160 mg of pure dry; el colesterol in el loro-form and transfer with wast ages of el loroform to a 100-ml flask. Vlake up to volume with elhoroform and mix Tl volume to solve the order of the standard transfer 5 ml of stock stan lard to a 100-ml volumetre drak dutte to the mark with elloroform and mix Tl is solution een tains 0.8 mg of el olesterol in 10 ml. It is stall be for some days if kej t. e. I i an l away from light.

cholesterol, in the same way, and for photometric measurement prepare a blank by treating similarly a 10-ml. portion of chloroform alone, After mixinc, keep in a dark place at 25° C, for exactly 30 minutes, 100 then transfer to colorimeter or photometer containers and read.

For colorimetric measurement, compare the unknown against the standard in the usual way. The colorimeter may he equipped with a light filter over the exeriece (Wratten No 71A or its equivalent, supplied by Fastman Kodak Co, or hy the manufacturer of the colorimeter) although this is not essential Since the standard color fades rapidly, if more than two or three unknowns are to be read, a number of standards must be prepared, suitably timed with respect to the unknowns, or the standard may he used to standardize an artificial standard (see previous method) scainst which the properly timed unknowns are read

For photometric measurement, determine the densities of the highly standard, and unknown at exactly 30 minutes after adding the acetic anhydride-sulfuric acid reagent, in a photometer at 660 mg. Set the photometer to zero density with water. Subtract the density of the blank from the other values to obtain their true densities

CALCULATION For colorimetric measurement. Since the 10 ml of extrict used correspond to 0 4 ml of original sample, the calculation is as follows

Reading of Standard $\times 0.8 \times \frac{100}{0.4}$ = mg eliolesterol per 100 ml

For photometric measurement

Density of Unknown $\times 0.8 \times \frac{100}{0.4} = \text{ing eholesterol in 100 ml}$

Under the conditions specified and in a 1-cm cuvette the density of the standard is approximately 0 700 Since the standard corresponds to a serum with 200 mg per cent cholesterol the limit of accurate measurement under these conditions is about 300 mg per cent It is suggested for photometric measurement that smaller aliquots such as 2 ml or 5 ml, of the chloroform extract be taken diluted to 10 ml with chloroform, and the color then developed as described. Using the above calculation formula results must then be multiplied by 5 or 2 respectively. In this way a much wider range of cholesterol concentration may be accurately covered

Discussion. The final color in the above procedure is due to both free cholesterol and cholesterol esters. Since free cholesterol and ester cholesterol give different amounts of color per milligram, and the color develop? at different rates, interpretation of results may be obscure if an abnormal distribution of cholesterol between free and ester forms is present. Reinhold 117 utilizes the difference in velocity of the Liebermann-Burchard reaction with cholesterol esters and free cholesterol as a basis for determination of cholesterol partition Sperry and Brandiis describe a procedure for total cholesterol after saponification of the ester fraction, thus eliminating errors duc to variation in ester content

The Liebermann-Burchard reaction with cholesterol gives much more color in chloroform than in the glacial acetic acid used in the previous method To offset this advantage, the reaction in chloroform is, according to Schoenheimer and Sperry, more sensitive to the effect of such analyti cal variables as time, temperature, etc. so that more careful technical

in Reinhold Proc Soc Expl. Biol Med 32 614 (1935) in Sperry and Brand J Biol Chem 150 351 (1943)

control is necessary. The spectrophotometric characteristics of the color in chloroform are somewhat similar to those of the color in acetic acid (see Fig. 153) except that peak light absorption in the red end of the spectrum is at 660 mm instead of at 625 mm 119 The high light absorption at 420 mu permits use of this range as well as the 660 mu range. Although the color intensity here slowly increases rather than waxing and waning as at 625 mu, and is a function of the extent of exposure to light during color development, neverthless Summerson and Rohmson 100 found this range to be in some respects more satisfactory than at 660 mu Sunderman and Razek 119 have shown that measurement at 530 mm is also satisfactory. although the optical density is low here, the agreement with Beer's law is excellent

Interpretation See under previous method

Other Methods Other methods for the determination of serum cholesterol usually employ isolation as the digitoride, followed either by weighing on a microbalance 101 or oxidation with chromic acid In the oxidation procedures either titrimetrie122 or gasometrie133 methods may he employed For details see the original papers

DETERMINATION OF FATTY ACIDS

Introduction. The fatty acids of blood are present largely as esters in the form of (a) neutral fat, (b) phospholipides, (c) cholesterol esters, there may be a small amount of free fatty and (i.e. soaps) also present The total fatty acids of whole blood are distributed approximately evenly between cells and plasma, this even distribution apparently simply re flects the fact that while phospholipides for example are much more abundant in the cells than in the plasma, the reverse is true for esterified eholesterol which is apparently absent from the cells. Total fatty acid content is usually determined by saponification and titration of the hherated fatty acids

Method of Stoddard and Drury 124 Principle The blood is extracted with alcohol-other the extract sanoussed the faits acids separated filtered washed dis solved in alcohol, and titrated with the mol blue as indicator

Procedure Extraction Five mi of whole blood, plasma, or serum is introduced gradually into a 100 ml volumetric flask containing about 75 ml of a mixture of 95 per cent alcohol and redistilled ether (3 1) The flask is Immersed in boiling water and rotated frequently and vigorously (to prevent superheating) until boiling begins, then cooled to room temperature, made up to volume with alcohol ether, mixed, and filtered through fat free filter paper

Saponification To a 100 ml beaker containing a few grains of coarse sand (previously boiled with acld, washed, dried, and extracted with ether) add

¹ Sunderman at i Razek J Biol Chem 118 3-9 (1937)

^{1 9} Upt ut lished

Unjutinsed
 Unjutinsed
 Mueller J Biol Chem
 St9 (1917) Ewert Bochem Z 263 159 (1933)
 Okey Proc Soe Expl Biol Wel 2c 518 (1999) Turrer J Biol Chem 92 495 (1931)
 Kirk Page and Yan Sike J Biol Chem 106 903 (1931)
 Stodlyr I and Drury J Bisl Chem 31 "41 (199) There et old has been critically exalured by Man and Gulder J Biol Chem 94 43 (1933)

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gradually, while evaporating on a water bath, 75 ml. of the filtrate. The temperature should be low enough to avoid perceptible boiling but may be raised after the ether has evaporated. Evaporate to a volume of about 30 ml. Add 0.1 ml, of saturated CO2-free NaOil, 125 mix, add a few grains more of sand, cover with a watch glass, and boil gently for 20 to 30 minutes (to saponify).

Separation and Washing. Remove the watch glass, drop in a small piece of litmus paper, make acid with 30 per cent liCi; then run back to alkalinity with 10 per cent NaOll (in order to avoid an excess of alkali on evaporating to dryness). Evaporate to dryness (in order to get rid of all alcohol). Add 15 ml. of water, heat on the steam bath, and stir to dissolve the soaps. While hot, add a drop of thymol blue indicator and make acid (faint pink) with 30 per cent 11Cl. Set the heaker for 10 minutes in cold water, then swirl almost continuously for five minutes to produce a better separation of the fatty aclds.

Filters are previously prepared as follows: Use a Gooch crucible, smallest size (top 28 mm., bottom 18 mm. in diameter). Set the crucible in a rubher holder which fits over a 500-ml, suction flask, A paper-pulp suspension is made by shaking up a piece of soft filter paper in 300 to 400 ml. of distilled water. Shake vigorously and immediately pour some into the crucible while there is a strong suction on. Repeat until a laver about 1 mm. thick is formed. Tamp the layer down carefully all over with the end of a glass rod. Allow the larger masses of filter pulp fibers in the suspension to settle out, and pour on successive amounts of the thin upper suspension of isolated shreds, keeping a strong suction on and tamping down occasionally, until the filter is dense enough to offer a definite resistance to the suction. Remove the crucible from the rubber holder and dry in an air oven at 110° C. for 15 minutes. Allow to cool before using.

Place the crucible in its holder in an ordinary funnel and filter some of the fatty acid suspension into a test tube. If the filtrate is not perfectly clear, put it through the crucible again. If the filtration does not start in a few minutes, transfer the crucible and rubber holder to the filter flask and start the suction very gently, with a test tube under the funnel. After filtration has started, continue without suction After the fatty acid suspension is filtered and drained, wash with 4 mi. of 5 per cent NaCi solution, neutralized to methyl red. Use a pipet and run the salt solution down the walls of the beaker all around; then, tipping the beaker, use a fine bent glass rod to rinse the side of the beaker more thoroughly with the solution, then pour this rinsing into the Gooch crucible, rinsing its side with the aid of the rod. Wash until the filtrate from one washing takes not more than 0.05 ml. of 0.02 N NaOll to neutralize it to phenolphthalein. Usually this is true of the third washing. Put the crucible back on the suction flask with a nonprotein nitrogen tube (cut off to a convenient height and calibrated at 1-ml. intervals from 10 to 15 ml.) under the funnel Wash down the walls of the beaker with 5 ml. of 95 per cent alcohol, heat to bolling, and pour into the crucible. With the glass rod quickly loosen up any fatty acid fragments on the wall of the crucible. Allow to run nearly out, then put on a moderate suction. Rinse out the beaker and crucible twice more with 3 ml. of alcohol each time, heating It to boiling Then wash off the outside of the crucible and the funnel.

Titration. Add a few grains of sand, boll the filtrate for 1 minute, cool in a beaker of water, note the volume of alcohol, add 3 drops of 0.3 per cent thy mol blue in 50 per cent alcohol, titrate with 0.02 N NaOl1 to a pure blue which stays practically unchanged (no yellow tinge) while shaking for 2

¹¹³ bee Appendix

minutes, keeping a stopper in the mouth of the tube to avoid absorption of GO: For a blank boil 10 ml of sicohol and titrate

CALCULATION Calculate the correction necessary for the amount of alcohol present before titration Add a correction amounting to 0 005 ml for each ml of NaOH used in titration (a simple correction for the volume of solution) Subtract the total correction from the titration Multiply by the normality factor thus getting the number of millimoles of fatty and Multiply by

of millimoles of latty acid. Multiply by $\frac{100}{\text{ml filtrate evaporated}} \times \frac{100}{\text{ml blood used}} = \text{millimoles fitty acid per 100 ml blood}$

To translate into terms of weight (not a very significant figure) multiply by an aver age factor for the fatty acids as they usually occur in blood = 2772. The molecular weights are so nearly alike that a coosiderable variation in the proportions will not affect the calculated weight by more than about 2 per cent.

Interpretation. Human whole blood has a total fatty acid content ranging from about 9 to 14 millimoles (milliequivalents) per liter This corresponds to about 250 to 390 mg of average fatty acids per 100 ml Some variation is found hetween different individuals, and in a particular individual at various times. Factors such as diet and disease which influence the blood content of neutral fat, phospholipide, and cholesterol esters will naturally influence the total fatty acid content

DETERMINATION OF LIPIDE PHOSPHORUS

Introduction. The lipide phosphorus of blood and tissues is found in such compounds as lecithin, cephalin, sphingomyelin phosphatidyl serine, etc, which are obtained by the extraction of tissues with certain nonaqueous solvents. In blood plasma, lecithin is the major phospholipide according to Bloor, while in the red cells cephalin and sphingomyelin predominate, the data of Kirk-125 present a somewhat different picture. The simplest procedure for the determination of phospholipide is the analysis of lipide-containing extracts for total phosphorus, as described below, this procedure gives fairly accurate results. Bloor 1 isolates phospholipides from nonlipide phosphorus by precipitation with acctone and magnesium chloride, and determines the phospholipide titimetrically after oxidation with chromic acid Gasometric determination as total carhon by wet combustion is also satisfactory. 128 Kirk-126 and Artom-126 have described procedures for the separate microdetermination of certain of the various phospholipides in blood.

Method of Youngburg, 12 Modified Principle The extracted lipides are oxidized with sulfure acid and hydrogen peroxide and the phosphate present determined colorimetrically. The phosphate procedure of Faske and SubbrRow is applied here the Youngburgs use the stanous chloride reagent of Author and Cohen (see p. 630) which is applieable to smaller amounts of phosphorus.

¹³⁶ Kirk J Biol Chem 123 623 637 (1938) See however Sinelair and Dolan J Biol Chem 142 659 (1942) Artom 65id 157 595 (1945)

[&]quot;Blood J Bod Chem 18 20 73 (1929) For an adaptation of this method see Ellis and Majnard J Biod Chem 18 210 (1937) McCos and Schultze U Biod Chem 18 479 (1941) describe a plotometric adaptation of the clorum acid oxidation procedure suitable

for small an ounts of h₁ ides

11 Airk Page and Van Slyke *J Biol Chem* 106 203 (1934) Van Slyke and Folch
ibid 135 509 (1910)

¹²⁰ Youngburg and Youngburg J Lab Clin Wed 16 158 (1930)

Procedure Transfer 18 ml of alcohol-ether mixture)** to a wide-mouthed test tube (hest 150 b; 20 mm) graduated at 20 ml, and drop in slowly, while shaking, 1 ml of plasma or serum. Mx, place in a boiling water bath, and heat the contents of the tube to boiling. Remove and allow to cool to room temperature. Make up to the 20-ml mark with alcohol-ether mixture, mlx, and filter.

and filter.

Transfer 8 ml of filtrate to a 200 hy 25 mm pyrex test tube, add a silica pebble (from broken silica ware), place in a wire rack containing a wire bot tom, over an electric hot plate, and evaporate to dry ness

Add 2.5 ml of 5.N sulfuric acid to the residue in the tube and digest over the hot plate as in the method for total acid-soluble phosphorus (see p. 633, including oxidation with perhydrol (30 per cent hydrogen peroxide). The remaining procedure is the same as for total acid-soluble phosphorus of blood (see p. 633), with color development at a final volume of 25 ml, a different standard is however used, containing only half as much inorganic phosphate, i.e. q.5 ml of standard phosphate solution, containing 0.04 ml of inorganic phosphate, instead of the 1 ml specified on p. 633. The could tions for colorimetric or photometric measurement of the color intensity are the same as for acid-soluble phosphorus.

CALCULATION Since the 8 ml of extract represent 0.4 ml of original sample calculation of results is similar to that for acid-oluble phosphorus except that the value 0.0 replaces 0.08 in the calculations corresponding to the use of half as strong a standard of calculate as for acid-oluble phosphorus and divide the result by 2 to obtain ms of lipide phosphorus per 100 ml of original sample

Interpretation. Plasma or serum contains about 9 to 10 mg of lipide phosphorus per 100 ml, whole blood slightly more, averaging about 12 mg per cent These values may be expressed as lecithin by multiplying by 25, since lecitlin contains approximately 4 per cent phosphorus, it is known, however, that only part of the total blood phospholipides is represented by legithin Artom (loc cit) gives the following distribution of phospholipides for a sample of human plasma lecitlin, 55 per cent of the total phospholipides, phosphatidyl ethanolamine (cephalin), 21 per cent, sphingomyclins, 12 per cent, pho-phatidyl serine, 7 per cent Little 15 known concerning the pathological significance of variation in this distribution Pathological variation in total phospholipide content is greater for red cells than for plasma In diabetes and nephritis, the lipide phophorus follows roughly the degree of hpema Increases in lecithm are also noted in pregnancy and in certain hepatic conditions. The existence of a constant ratio between cholesterol (which is antihemolytic) and lecithin (hemolytic) has not been definitely established

DETERMINATION OF BILE PIGMENT IN SERUM

1 Icteric Index (Meulengracht) Principle The intensity of yellow pigments tion of serum is compared with a standard pota-sum bichromate solution

Procedure ¹³¹ Separate the serum from 4 or 5 ml of freshly drawn unhemo lyzed blood For colorimetric measurement, accurately dliute i ml of serum

¹²⁴ Alcohol Ether Murture Three volumes of 95 per cent redistilled alcohol and 1 volumes of redistilled effer Mix.

¹³¹ Reagents Required 6 Per Cent Softum Citrate Solution Dissolve 50 g of USP of CP sodium citrate in water and dilute to I liter. If turind, allow to stand several days and

with 5 per cent sodium citrote solution in a small graduated cylinder, according to its color, until an opproximate match with the standord potas sium bichromate solution is obtained Transfer to a colorimeter cup and compare account the standard bichromate

For photometric measurement, dilute 1 ml of serum to 10 ml with 5 per cent sodium citrate solution. Determine the density in a photometer at 420 m μ_s setting the photometer to zero density with the 5 per cent sodium citrate solution alone. Determine the density of the standard potossium bichromate solution at the same wovelength, using water as a blank

CALCULATION For colorimetric measurement

 $\frac{\text{Reading of Standard}}{\text{Reading of Unknown}} \times \text{Dilution} = \text{Icteric Index}$

For example if 1 ml of scrum is diluted to 6 ml the dilution is 6 and if the unknown exactly matches the standard in the colorimeter the actions index is 6 0

For photometric measurement. Since the standard bichromate solution has an equivalent reterie index of 10 under the conditions of the procedure, the calculation is as follows.

Density of Unknown Density of Standard × 10 = Icteric Index

In a 1-cm cut ette at 420 ms an seteric index of 10 gives a density reading of ap proximately 0 200 at a 1 10 dulution of the serum. Under these conditions the limit of accurate measurement corresponds to an interior index of approximately 50 For higher values the determination is repected at a greater dulution of the serum, the own dulution value replicing the 10 in the obove calculation formula. Since the stand and is a stable colored solution is decostly should be constant. Therefore once the density of the standard has been estoblished for a given photometer and wavelength in filter, this value may be used in the future without the accessity of repeating the reading of the strandard in every series of onallyses.

Interpretation. The normal leteric index ranges between 4 and 6 The zone of latent jaundice (i.e., hyperbilirubinemia without elinical signs of jaundice) is between 6 and 15 Above this value interior symptoms may be observed. The yellow color is considered to be due chiefly to the presence of bultrubin per 100 ml of serum. The presence of certain other pigments will lead to errors, hemolysis in particular is to be avoided. This may be done by using a dry synnge and needle for collecting the blood, allowing the blood to clot protected from light, and centrifuging sharply to obtain a clear serum. Blood should be drawn before breakfast to avoid chyle, hipemic sera cannot be used because of interfering turbidity. The carotinemia which follows ingestion of carrots will lead to a high apparent letteric index and false interpretation, carrots should not be caten the day before the test. Further aspects of the significance of serim bilirubin are discussed below.

filter Add a little chloroform as a preservative Saline solution (0.9 per cent sodium chlor ide) may also be used as a diluent in the colorimetric procedure—but citrate gives clearer solutions.

Nan lard Potass: m Bichromate (DOI Per Gent). Dissolve exactly 0.1 g of reagent grade potassaum bethromaten water transfer with manage to all liter volumetre flash, and 12 to 3 drops of concentrated sulfure and dilute with water to the mark, and mix. Keep in a brown glass pottle. The exterie index of this solution is arbitrarily defined as equal to 1

CALCULATION Density of Unknown \times ing bihrubin in standard \times $\frac{10}{1} \times 100$ Density of Standard

= mg bifirubin p∈r 100 ml scrum or plasma

The density of the standard is estal lished as follows. Transfer 4 inl. of dilute standard alcoholic solution of bilirubia contaming 0.03 mg of bilirubia to a test tube and add 5 ml of methyl alcohol followed by 1 ml of diazo reagent. Mix by inversion and allow to stand 30 minutes for color development. At the same time, prepare a black tube containing 3 ml of methyl alcohol and 1 ml of diano reagent. Mix. Micr 20 minutes, set the photometer to zero density at 540 mm with the blank tube and deter mine the density of the standard

Under the conditions specified and in a t-cm curette the standard has a density of approximately 0 300 (see Fig. 156). Since this standard corresponds in an analysis to a

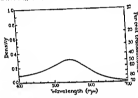


FIG 156 ABSORPTION SPECTPEM OF AZO-BILIRURIN, AS OBTAINED IN MALLOT-EVELTY BILIBURES METHOD

serum bilirubin content of 10 mg per cent up to about 30 mg per cent of bilirubin may be accurately determined. If the value is above this dilute the contents of both blank and unknown tubes with an equal volume of 50 per cent methyl alcohol, read immediately, and multiply the results by 2

The color developed in this procedure is quite stable and reproducible Since bilirubin is expensive and the standards do not keep very well it is feasible to determine the standard density once carefully for a given photometer and wavelength setting of filter, and to use this value in future analyses without the necessity of repeating the reading of the standard each time

If a photometer is not available the unknown may be compared in a colorimeter against the standard in the usual manner. A green filter over the eyepiece of the colorimeter will aid in color matching. An artificial standard containing cobalt sulfate bas been suggested for visual colorimetry," to replace the bilirubin standard. A colorimetric method has been described by Gibson and Goodrich 145

Interpretation. Normal plasma or serum contains 0 1 to 0 25 mg of bilirubin per 100 ml Bilirubin content is sometimes expressed in "units," 1 unit corresponding to 05 mg per cent, thus normal plasma contains 02 to 05 unit The zone of latent jaundice is represented by the range of 0 25 to 10 mg per cent Hyperbilirubinemia may occur in diseases of the liver or biliary tract, and also in extrahepatic conditions of a hemolytic

¹⁴⁴ White Brut J Expil Path 13 76 (1932) 145 Gibson and Goodrich Proc Soc Expli Biol Med 31, 413 (1934)

nature, such as those accompanying infectious diseases, permicious anemia, hemolytic anemia, hemorrhage, etc Low values for bilirubin may be found in secondary anemia. The determination may therefore be of importance in the differential diagnosis of anemias. The renal threshold for bilirubin is 3.5 to 4.0 units (1.7 to 2.0 mg per cent) in the plasma.

THYMOL TURBIDITY AND FLOCCULATION TEST

Principle Maclagan¹⁴ discovered that thymol, in barbital buffer at low ionic strength, added to serum produced marked turbidity in the presence of prenchymatous liver disease. In addition, doculation often appeared on longer standing

The thymol barbital reagent is saturated with thymol Temperature and other factors influencing solubility must be considered in preparing the solution. The procedure described 100 permits better control of thymol concentration and pH and yields a more uniform reagent of improved stability.

Procedure 14 Measure 6 ml of thy moi barbital reagent into a photometer cuvette. Place the tubes in a water bath maintained at 25° C \pm 1°. Add 0 1

lattle CO; it too alkaine Expured arr may be used as a consement source of CO;
The optimal temperature at which to keep the reagent appears to be 15° however at a
room temperature of 25° C the reagent remans unchanged for at least two weeks. At
temperatures lower than 15° it is more stable but crystals max separate thus causing lower
values for thymol turbidity. Remore sufficient solution for one day a use and place it in a
bath at 25° C at least 30 manutes before thing Exposure to carbon dioxide should be kept

to a minimum. The reagent should be renewed when it becomes opalescent

Standards Turbuitty can be measured either visually or photometrically. The latter is recommended For photometric measurement colloidal glass suspensions may be used as standards. These may be obtained commercially or prepared according to the following procedure. Fill a glass-stoppiered py rex reagent bottle of 1 liter capacity to about one-fourth its capacity with fragments of clean pyrex glass. Cover the glass fragments with distilled water and agitate vigorously on a mechanical shaker until a milky suspension is produced 8 to 20 hours its ordinarily sufficient. Decant the suspended glass into a 1000 ml cytuder and dilute to the mark with distilled water. Mix and allow to stand 14 days. Decant the upper 500 ml for use as a stock standard.

Standardization. I or this operation a Beckman Model DU spectrophotometer is satisfactory other instruments of comparable quality, should serve as well. Make a trial reading of stock standard in the spectroplotometer at a wavelength of 660 mg in a 10-mm cuvette using wider as a blank. Add sufficient water to an aliquot of the stock standard to make the absorbancy of the mixed diluted suspension approximately 0.100. Now measure the exact absorbancy of the diluted suspension. In absorbancy of 0.100 has been found to be equivalent to 0.30 Michagan units and 0.55 mm in longiand units in the 1 vely in plotoelectric colorimeter with a 0.00 mm filter 0 mil aperture and stan lard reflector. Thus in the cicladization of results, where 0 = 0.100 C = 3.3 (Machagani of 0.5 (Shank Hongland). It's is slightly above or below 0.100 the appropriate value for C may be calculated by direct proportion bor colorimeters of er than the 1 vely in instrument it is advisable to standard

¹⁴⁶ Maclagan Brit J Exp Path 25 234 (1944)

¹⁴⁷ Reinhold and associates Personal communication

¹¹⁸ Reagents Required Thymol Barbatal Reagent Transfer 60 g of thymol erystals (colories, b) to a 2000-ml pyrex Felemeser flask. Heat 1000 ml of datilled water to boding in another flask and bod it for about five minutes to remove carbon dioxide. Allon the water to cool to about 93° and pour about 300 ml into the flask on the thymol which will melt and partially dissolve. Add 309 g barhatal and 169 g sodium harhatal and theremander of the hot water to the flask containing the thirself without delay stopper the flask and mix by rotating vigorously for about five minutes Allow the solution to cool gradually to room temperature. Transfer the reagent to a volumetric flask of 1000 ml capacity and dilute to the mark with distilled water. Return the solution to the original flask. Idd about 1 g of thymol crystals. Shake vigorously until the solution becomes clear. Allow the flask to stand at 25 ± 1° overnight preferably in a constant temperature that hix again and filter dirough Whitman No I paper maintaining the temperature of the solution at or near 25° C. Test the pH which should be 7.55 ± 0.03. The pH may be adjusted by a dading 0.1 N. "Acil 11 the respect to the solution at or near 25° C. Test the pH which should be 7.55 ± 0.03. The pH may be

2. Van den Bergh Test ¹¹¹ Principle. Plasma or serum is treated with l'Inhibit resgent (dirzotized sulf indic acid). Bilimbin present riacts with the reagent to form a colored compound known as a varil in or a of diribbin ¹¹² Quishtatively, the color may develop immediately (direct rivation) or on standing ("indirect reaction) & casionally an atypical color divelops almost immediately). ("Implicate reaction") Quantitatively, the color may be used as a measure of the bilimbin content of the serum or plasma (see Section 3, p. 53).

Procedure 114 Obtain unhemolyzed plasma or serum as described abort Jaundiced serum must be diluted 1 5 or, if highly colored, 1 10, with water To 1 ml of serum (generally diluted), add 2 ml of freshly prepared Fhilchs diazo reagent 115 Direct reaction a pink to purple color develops immediately, reaching maximum intensity in a minute or so Indirect reaction acolor change is noted during the first 2 minutes. Within 10 minutes a golden color forms, changing over a period of an hour or more through brown to pink 115 Biphaic reaction a color appears during the first 2 minutes, but it is brownish red rather than the typical pink or purple of the direct reaction

Interpretation. The chemical and clinical significance of the three types of Van den Bergh reaction has been the object of considerable in vestigation Van den Bergh beheved that the different reactions were dependent upon the path by which the pigment cutered the serum For example, the direct reaction is usually found in the condition of obstruc tive jaundico, i.e. when the hihary passages are blocked, preventing the excretion of bile pigment which has already been secreted by the liver cells The indirect reaction is usually found when the jaundice is of hemo lytic or extraliepatic origin, and the hiphasic reaction is thought to represent a combination of both types Turther study has not justified the belief that the various types of launchee may be distinguished in terms of the qualitative Van den Bergh test 137 While normal sera and sera from cases of hemolytic jaundice usually show no direct-reacting bilirubin in most other types of jaundice it is usually possible to demonstrate the presence of both direct-reacting and indirect-reacting bilirubin It ap pears probable that there are two distinct types of serum bihrubin, for

uii Van den Bergh Presse méd 27 441 (1921) Sepulveda and Osterberg J Lab Clin Med 28 1359 (1943) See also Ducci and Watson J Lab Clin Med 30 293 (1945)

in proceedings of the process of the

In Reagents Required Dirac Reagent—Squiston A To 1 g of sulfainlin acid add 15 ml of concentrated hydrochloric send Add water stir to dissolve and dilute to 1 liter with water stir to dissolve and dilute to 10 liter with water stir to dissolve and dilute to 100 ml Just before the test prepare the distor reagent by mixing 25 ml of Solution A with 0.75 ml of Solution B. This mixed solution to the solution and the solution of the solution and the solution and the solution to t

This mixed solution keeps only a short time at room temperature

11 Better results are sometimes obtained by adding the diago reagent slowly down the
side of the test tube containing it a serum so that a layer is formed over it e serum Observe
the color at it e interface between serum and reagent

in The indirect test is fre mently entried out by adding three to four velumes of alcohol to the mixture of strom an I reagent after 10 mm ites. The indirect color appears immediately after inning.

ii See Greene 1 lotz an l Localio Arch Int Wel 61 658 (1938) Sepulveda and Oster berg 914 72 372 (1943) also pp. 402 404 in Bo lansky and Bo lansky Bochemistry of Disease and ed New York The Mactualian Co 1952

instance, indirect-reacting bilirubin may be removed from serum by extraction with chloroform, while direct-reacting bilirubin is not soluble in this solvent Griffiths138 has reported the isolation from human bladder bile of a pigment called "cholebilirubin" which gives the direct reaction. is distinct chemically from bibrubin, and appears to have the properties of the direct-reacting bilirubin of serum. It has been proposed that the terms "cholebihrubin" and "hemobilirubin" be used for the directreacting and indirect-reacting bilirubin respectively, these terms carry the unfortunate implication of origin, which has not been conclusively established Other interpretations of the Van den Bergh reaction which have been proposed include the theory that variations are due to concentration differences. 129 or that the bilirubin of plasma is bound to protein. with a portion capable of dissociating to give an immediate reaction ("direct-reacting") while the remainder ("indirect-reacting") can react only slowly or after the addition of alcohol 140 No single theory has apparently found universal acceptance 141

3 Quantitative Determination of Serum Bilirubin (Method of Malloy and Evelyn) 141 Principle The principle is the same as that used in the qualitative Van den Bergh reactions described above. The "azobihrubin formed is determined by photometrie measurement

Procedure 143 Dilute 1 ml of unhemolyzed plasma or serum to 10 ml with water, and mix Blank Place 5 ml of absolute methyl alcohol in a test tube Add 1 ml of the hiank hydrochloric acid solution, mix by tapping, and add 4 ml of the 1 10 diluted serum or plasma Mix gently by inversion Unknown Place 5 ml of absolute methyl alcohol in a second test tuhe Add 1 ml of freshly prepared diazo reagent, mix by tapping, and add 4 ml of the i 10 diluted serum or plasma Mix gently by inversion. Care must be taken to handle both blank and unknown tubes in the same manner, so that any turbidity which may form will be the same in both tubes. Allow to stand 30 minutes for color development, then transfer to suitable containers and read in the photometer at 540 mg. Set the photometer to zero density with the blank This compensates for any extraneous color or turbidity present in the unknown

¹¹⁴ Griffit1 4 Biochem J 26 1155 (1932) See also Najjar Pediatrics 10 1 (1952) Colo and Little J Clin Path 6 99 (1953)
115 Sinder and Rienhold Am J Med See 130 248 (1930)
114 Cololige J Biol Chem 133 113 (1940) Vlartin J Am Chem See 71 1230 (1949)
115 For review see Gray The Biol Pymaries New 1 ord John Wiley & Sons 1953
116 Nailey and Lively a J Biol Chem 194 431 (1937)
117 Marie Peron Service and The Lee of suffernite seed add 15 ml of concentrated hydro-

Diazo Reagent Solution A To 1 g of sulfamilie acid add 15 ml of concentrated hydrochloric acid dissolve and dilute with water to I liter Stable indefinitely Sol tion B Dis solve 0.5 g of sodium mitrite in water and dilute to 100 ml keep away from hight and discard when it turns I ellow Dia o Reagent Mix 10 ml of Solution 1 with 0 3 ml of Solution B Must be prepared fresh before using

Blank Hydrochloric Acid Solution Dil ite 15 ml of concentrated by drochloric acid to 1 liter with water Stable indefinitely

Standard Bilirubin Sol tion Stock Standard Place 10 mg of bilirubin (obtainable from Lastman Kodak Co Nochester VV or Hoffma in I a Rocke Vutley VJ) in a 100-int volumetric flash add el loroform to dissolve and dilute to 100 ml with chloroform Keep in volumetric hask and ci foroiorii to use at lard. Transfer 10 ml. of stock standard to a 100-ml. volumetric flask and d lute to 11 e mark with methal alcohol. Mix. This standard contains 0 01 mg of librubin per ml Prepare just before using

596 ml of serum from a pipet that will deliver accurately hetween marks (Do not blow out the pipet) Stopper the cuvette and mix well in 30 minutes measure the absorbancy (optical density, see p 5t3) at 660 mm, using thy mol barbital solution for the zero setting. If the thymol turbidity exceeds 20 units (see below), repeat using 0 05 mt of serum Multiply the result by 2 when this is done

nen this is none Calculation. Thymol turbidity units $=U imes rac{C}{5}$ where U is the absorbancy of the unknown S of the glass standard and C is the thymol turbidity equivalent of the etendard

For visual measurement, the egg allumin gelatin formazine standards of kingsbury et al developed for the determination of protein in urine (see p 929) were originally used by Maclagan Companson against these standards should be made with light coming from helind the observer Calculate results by dividing the protein equivalent of the standard most closely matching the unknown by 60 to give Maclagan anits and by 30 to give Shank Hoaglan I units

Thymol Flocculation Decant the turbid solution into a conical centrifuge tube of 15 ml capacity Stopper and replace in the water bath at 25° C until the following day Framine the solution for the presence of a flocculum and note also the extent to which the supernatant fluid has cleared Flocculation is graded on a scale from 4+ (water clear) to 1+ (flocculation and clearing are minimal)

Interpretation: a Thymol Tunnibity Results may be expressed either as Maclagan units or as Shauk-Hoagland units One Maclagan unit equals two Shank Hoagland units The Commission on Liver Discase of the Armed Forces Epidemiological Board has recommended that Shank-Hoagland units be adopted in preference to Maelagan units Minety-five per cent of a group of healthy subjects tested by Reinhold and associates were found to have thy mol turbidities not exceeding 50 Shank-Hoagland units, and 99 per cent were below 6 6 units, using the procedure described here

The thymol turbidity test is among the tests most frequently positive in virus hepatitis, and is particularly useful during the recovery period for evaluation of progress and for detection of carriers of viral hepatitis Although patients with eirrhosis may show elevated thymol turbidity, it may fail to become positive in cirrhosis either of the Laennec or biliary

The thymol turbidity value is consistently within the limits of normal in biliary obstruction of recent origin older lesions may occasionally result in abnormal values Since the reagent also reacts with serum lipides blood should be obtained from the patient in the postab-orptive state, and confirmatory evidence sought when serum lipides are elevated Thymol turbidity may also be increased in any disease characterized by an elevated serum \(\gamma \)-globulin content quite independently of liver involvement such conditions include multiple myeloma lymphogranu loma sarcoidosis parasitie infections and others. While the liver may also be involved proof can only be obtained by recourse to other liver function tests not directly dependent upon changes in serum protein content

h THYMOL FLOCCULATION Climeal experience suggests that the occurrence of flocculation depends upon some additional or different factors than does the production of turbidity 149 In general, the significance of a positive flocculation test is the same as for ahnormal turbidity The flocculation test is less frequently positive than is the turbidity test. however, false positive tests are uncommon and the occurrence of thymol flocculation thus may he accepted with a higher degree of confidence as evidence of liver damage. The serum of healthy individuals shows no flocculation Flocculation graded 1+ or more, therefore, is abnormal

CEPHALIN-CHOLESTEROL FLOCCULATION TEST

Principle Suitably diluted serum of patients suffering from liver disease forms a flocculent precipitate when treated with a suspension of cephalin and cholesterol in water 150 The mechanism of the reaction differs somewhat from that of the thymol turbidity test, the change in serum responsible for the test appears to be primarily in the albumin fraction 151

Procedure 132 It is advantageous to use conical centrifuge tubes of 15 ml capacity Measure into such a tube 4 ml of 0 85 per cent sodium chloride solution Add 0 2 mi of serum and mix well by tapping the tube vigorously 5 times Add i ml of cephalin-cholesterol reagent and again mix by tapping the tube vigorously at least 15 times Place the tube in a water bath at approximately 25° C, in the dark and away from chemical fumes, until the following day Examine the tube for presence of flocculation after 24 hours and again after 48 hours (optional)

Grade the reaction as 1+ to 4+ as described under "Thymoi Flocculation" (p 596) taking into account the amount of precipitate and the transparency of the supernatant fluid If desired, the precipitate can be removed by centrifugation, the supernatant fluid decapted, and its absorbancy measured in a

¹⁴ Neefe Gastroenterology 7, 1 (1946)

 ¹¹⁰ Hanger J Chn Invest 18 261 (1939)
 151 Moore Pierson Hanger and Moore J Clin Invest 27 737 (1948)

Moore Pierson Hanger and Moore - Clin ances A. The purchase of the reagent is Reagents Required Cephalin Cholesterol Stock Solution The purchase of the reagent (antigen') is recommended (Difco and Wilson brands are satisfactors). It is supplied in vials containing 100 mg of cephalm prepared from bram and aged and 300 mg of choles terol The contents of a vial are dissolved according to directions in USP ethyl ether after addition of 1 drop of distilled water. The stock solution so obtained may be kept for

several months if refrigerated and tightly atoppered Cephalin Cholesterol Reagent Measure 32 ml of water into an Erlenmeyer flask of 50 ml capacity with a mark scratched on it to indicate 27 ml. Place the flash on a water bath on an electric heater and heat to 65° or 70° C While the water in the flask is kept at this tempera ture add drop by drop I ml of the stock cephalin-cholesterol solution rotating the flask at the same time Place the flask directly on the hot plate at low heat and boil gently to remove the ether until the volume is reduced to 27 ml When cool add 3 ml of merthiolate solution aqueous 1 1000 as a preservative This reagent retains its activity unchanged for at least a week if stored in a refrigerator

Precautions Preparations of the cephalin-cholesterol reagent as purchased or prepared have been found to vary wilely in sensitivity. Fach lot of reagent must be assayed by testing serum of healthy individuals and of a sampling of patients showing positive reactions Acceptal le reagents do not produce more than an occasional 1+ flocculation when the sera of lealtly subjects are tested and on the other hand yield a high proportion of positive ficeculations in patients suffering from liver disease. Once a sintable reagent has been obtained new lots can be tested by comparing them with previous ones for best results tests aloud be done in duplicate using two different preparations. Difficulties of standardizing and of controlling this test make it practical only when a considerable flow of samples through the laborators may be expected

arm by means of a different springe and needle. The exact time of completing the injection and of collecting the blood should he noted. The blood is allowed to clot, and serum is removed for analysis. Care is used to avoid hemolysis. Needle and syringe should be dry, and the blood should be transferred from syringe to tube gently and without production of foam.

MEASUREMENT OF DTE CONCENTRATION Measure 0.5 ml of the serum of the dyed specimen into a cuvette containing 5.0 ml of water. Add 0.5 ml of 0.5 N sodium hydrotide solution Mis well and measure the absorbancy (optical density) at 6.60, 555, and 420 mg, using water for the zero settlings at each wavelength Maximum absorption occurs at 550 mg and this wateringth settling is theoretically to be preferred, however, interference from hemoglobin is less marked at 565 mg if Serum billirubin is elevated, the control sample is diluted and absorbancy measured as described for the dyed sample. The figures so obtained are subtracted from those of the brom sulfatein containing serum. Calculate results as described helow.

CALCULATION Bromsulfalein mg per $100 \, \mathrm{ml} = f \times (U_{44} - 1.28 \, U_{44}) = 0.15 \times (U_{45} - 1.28 \, U_{45}) = 0.15 \times (U_{45} - 1.28 \, U_{4$

$$f = \frac{C}{S_{ext}}$$

where C is the concentration of standard and S₄₄ is the absorbancy Average the values to give the value of f used in the calculation. Once f has been determined standards can be dispensed with except for occasional instrument checks. From the brompulishen content of the scrum the per cent retention is calculated as follows.

Per cent retention = bromsullalein in mg per 100 ml × 10

A correction is applied in the method described to compensate for the presence of hemoglobin in serum since hemoglass of some crythrocytes may be difficult to a rod and i-moglob in will cause falsely high readings even at 550 m_p. The effect of latesttence in serum is climinated by making the absorbancy measurement at 650 ms, where neither i romsulfatem nor hemoglobin absorb light. A correction may be used when the test is applied to jaundered patients.¹⁹⁸

Rownthal and White used 2 mg of dye per kg of body weight. The 5 mg per kg dosage however I as been adopted widely and provides a more sensitive and more

¹¹⁷ Ti e correction factors given in the calculation were calculated from readings with an Evelyn pl otocolorimeter (see p 526) They may be applied to readings made with other phot meters provided the transmittancies at the wavelength's used do not differ greatly from those obtained with the Evelyn instrument. If desired the factors may be determined by measuring the absorbancy A of a ddute solution of hemoglobin containing roughly 10 to 20 mg hemogloi in per 100 ml at 420 and 505 mg. The correction for hemolysis is calculate i by means of the ratio Ass./Ass obtained from the readings of the hemoglobin solu tion This corresponds to the factor 0 15 in the calculation. The turbidity correction is based on readings of a colloi al glass suspension similar to that described under thymol turbid ty (see p 595) made at 420 505 and 660 mm. Although turbid solutions have no absorption bands their absorbanes increases progressively from the red to the blue regions of the spectrum Therefore the ratio Asss/Asss of the glass suspension represents the factor t) which any measurements made at 660 mm must be multiplied and Acro/Aco the cor responding factor for the reading at 420 main order to correct for turbidity. In the calcula tion above these ratios are t 28 and 1 95 respectively. The turbility correction will vary with particle size and therefore represents an approximation. No corrections need be applied to the standards 112 Zieve Hanson and Hill J Lel Chn Med 37 40 (19.11)

precise test. The existence of an enteroheritic circulation of bromsulfalein in does not appear to influence the results sufficiently to necessitate a return to the original dosage 100 The time at which the postinjection blood specimen is collected is not of great importance provided that appropriate normal standards for that time are used the 45-minute sampling time has proved to be satisfactory.

Interpretation. Healthy individuals, after injection of 5 mg bromsulfalen per kg of body weight, retain less than 10 per cent at 30 minutes and 7 0 per cent at 45 minutes At 60 minutes, no dye is retained

Bromsulfalem retention is generally accepted as the most sensitive and dependable among the laboratory procedures currently used to demonstrate involvement of the liver It is especially helpful for evaluating suspicious or positive results obtained by means of floculation tests in the absence of byperbilirubinema. The bromsulfalem test outranks all others in the proportion of positive tests found in Laennec's cirrhoss. It has been among the most useful for following recovery from virus hepatitis and for detecting residual liver damage from this disease. It is probably the only procedure capable of detecting fatty liver, although it cannot be depended upon to do so consistently.

For the study of the naundiced patient the bromsulfalein test has hittle to offer Maximal retention occurs in the presence of severe liver damage, and further deterioration of the liver function can have no additional effect on die retention. Thus it is rarely used when hyperbili-

rubinemia or clinical jaundice are present

DETERMINATION OF PLASMA PROTEINS

Introduction. The standard method for the determination of protem is by the Kjeldahl type of digestion and ordation converting the introgen present to the form of ammona, which is then determined If the material contains nonprotein introgen, this is determined in a separate analysis and subtracted from the total introgen value to give the protein introgen. This is then multiplied by 6.25 to give the protein value since the average protein contains 16 per cent introgen. If sufficient material is available the most accurate version of this procedure is a micro-Kjeldahl method similar to that described in Chipter 31 for the determination of the total introgen of the inner For blood analysis with its inherent limitation in the amount of material available this is replaced by the various micro-Kjeldahl methods and the procedures become similar to the determination of nonprotein introgen already described (p. 545).

To obtain satisfactory results more rapidly and simply than is possible by micro-Lyeldahl methods various direct colorimetric, turbidimetric, and specific gravity procedures have been developed. The colorimetric procedures nre usually based on color reactions specific for protein or constituent amino acids and are standardized necordingly. Turbidimetric methods rely on the comparison of the turbidity produced by precipitating reagents on proteins in dilute solution with the turbidity produced similarly on a standard protein solution. The specific gravity methods

¹¹ Lorber at 1 St av Castroe sterology 20 20' (1952)

¹⁰ Owen J Lab Clin Ved 38 583 (1951)

are based on the fact that protein is by far the most abundant solid constituent of blood, and hence the specific gravity should be determined largely by the protein content

Tractionation of the plasma proteins prior to separate analysis is based almost entirely on the classical methods using concentrated salt solutions and the bulk of climical literature is in terms of such fractions. Fractionation by electrophoretic mobility has not yet reached the stage of routine practicality, except possibly in a few of the larger institutions. For a further discussion of the various methods of fractionating the plasma proteins see Chapter 22. The determination of blood hemoglobin content will be considered in a separate section.

1 Micro Kyeldahl Method ¹⁴ Principle Total proteins are determined in serum or plasma by a micro-Kyeldahl method employing direct nesslerization making the appropriate correction for nonprotein mitogen Fibrinogen in plasma is determined by solution as fibrin followed by digestion and direct nesslerization. Albumin is determined by analysis of the fluid remaining after precipitating the globulin fractions of 23 per cent sodium sulfate solution. Globulin is serum is estimated by subtracting the albumin from the total protein content in plasma by subtracting albumin and fibrinogen from total protein.

Procedure ¹² Total Proteins (Albumin + Globulin + Fibrinogen) Dilute es actly 1 ml of the serum or plasma to 50 ml in a volumetric flask with 69 per cent sodium chloride solution (If serum is used, the "total protein" will

in T) is method is a comb ned adaptation of the procedures of various authors. The direct nesolerization in cro-hyddahl procedure of Wong (J. Biol. Chem. 55.427 (1923)) employing perulifate for oundation in secommended because of its superiority when proteins is prient. Fractionation of the proteins with sodium auditate solut on follows the directions of the proteins with sodium auditate solut on follows the direction of the proteins with sodium auditate solut on follows the directions are superiority for the use of effect of aid in separating the albumin and globulin fractions as suggested by Kingeley Culton and than Shike (J. Biol. Chem. at 1.557 (1920)). These latter authority according to extra the superiority of tuning a uniform amount of anticoaculant such as oxialtate in comparative such so make the superiority of tuning a uniform amount of anticoaculant such as oxialtate in comparative such so make the superiority of tuning a tractionation section section and alter on variet attribution. Every cells and į lasma. They use 5 mg of potassuum oxialate per ml of blood. For other method of digestion and ifractionation sec Campbelli and Hanna J. Biol Chem. 118 19 15 (1937). Fillener and illuted insom 3°11.185.299 (1945). These latter authors use cold methyl above.

represent only albumin + glnbulin.) Using 1 ml. of this diluted mixture, proceed as directed below under "Digestion."

Fibringen. To 1 ml. of plasma in a small cylinder add 30 ml. of 0.9 per cent sodium chloride solution, folinwed by 1 ml. nf 2.5 per cent calcium chloride solution. Mix with a slender pointed glass rod, leaving the rod in the mixture. Allow to stand for 30 minutes or until a solld clot has formed, Carefully rotate the rod in the jelly, squeezing out the water hy pressing the clot against the side of the cylinder at the same time. All the fibrin should stick to the rod and appear ultimately as a thin white sheath over the rod. If any hits of clot escape this process, pour the cylinder contents onto a dry filter. and when the fiuld has drained pick up the remaining clots with the tip of the rod and squeeze out the excess fluid by pressing against the side of the funnel. Transfer the rod and adhering fibrin to a piece of dry filter paper and dry as thoroughly as possible by rolling against the filter paper. Transfer the rod and dried fibrin to a centrifuce tube graduated at 10 mi, and containing 4 ml, of 1 per cent sodium hydroxlde solution. Place tube and contents in a boiling water bath until the fibrin has dissolved and only a turbid suspension of calclum oxalate remains. This should require but a few minutes. Remove from the water bath, remove the rod, washing it down with a few ml, of water, and make up the contents of the tube to 10 ml, with water. Mix and centrifuge. Pipet a 5-ml, portion of the clear supernatant fluid to a digestion tube and proceed as described helow under "Digestion."

Albumin. To 1 ml. of serum or plasma in a 59-ml. centrifuge tube add exactly 30 ml. of 23 per cent sodium sulfate solution. Stopper and mlx by Inversion. Add about 5 to 10 ml. of ether, again stopper, and shake vigorously. Centrifuge for about 10 minutes, capping the tube to prevent loss of ether. After centrifuging, the precipitated globulins should form a compact layer below the ether and above the clear albumin solution. Siant the tube and lasert a pipet of narrow bore, and with the mouthpiece closed by the finger, along the side of the tube past the packed globulin layer into the clear fluid below. Fill the pipet with the fluid and transfer to a dry test tube, wiping off any precipitate adhering to the nutside of the pipet before discharging its contents. Use 1 ml. of this for digestion as described below. If a centrifuge is not available, the mixture may be poured onto a retentive filter (such as Whatman No. 50) and covered with a watch glass to prevent evaporation. If the first portions of filtrate are not clear, return to the filter. Use 1 ml. of the clear filtrate.

Digestion. Into a pyres test tube graduated at 35 and 50 ml. (see method for nonprotein nitrogen, p. 545) place 1 ml. (or 5 ml. in the case of fibrinogen) of the solution to be analyzed. Add 1 ml. of 1:1 sulfuric acid and a quartz chip or a few glass beads. Digest over n microburner as described for the determination of nonprotein nitrogen until excess water has been driven off, the solution darkens, and white fumes appear. When the tube is nearly full of dense fumes, cover the mouth in the tube with a watch glass and reduce the flame or raise the tube an that the mixture boils gently. Continue boiling for 3 minutes. Remove the burner and allow to cool for 1 minute. Add to the tube contents, drop by drop, 0.5 ml. of persuifate solution. Replace the burner, tap the tube to start boiling if necessary, and con-

¹⁴ On cold days some of the sodium sulfate may crystallize out during the centrifuging. If this happens warm the tube and contents in the mulbitor at 37° C until the crystals have redissolved, add more ether, shake and again centrifuge as described.

tinue holling until clear Cool, dilute with water, and proceed with the direct nesslerization and colorimetric or photometric measurement exactly as de scribed on p 516 for the determination of nonprotein nitrogen. The same standard is used, containing 0 15 mg of nitrogen, in the presence of 1 ml of the 1 I sulfuric acid and 0 5 ml of persulfate solution, to balance the amounts present in the unknown For photometric measurement the blank tube cootains water, acid, and persulfate as described for the standard.

CALCULATION For colorimetric measurement Total Protein and Albamin are calculated directly as follows

Total Protein and Albamin are calculated directly as follows:
$$\left[\left(\frac{\text{Reading of Standard}}{\text{Reading of Luknown}} \times 0.15 \times \frac{100}{1} \right) - \text{NP} \right] \times \frac{\text{C 25}}{1000}$$

= grams protein per 100 ml (or per cent protein)

where I represents the actual volume of serum or plasma used in the determination NPN represents the nonprotein nitrogen content in mg per cent as determined in a separate analysis. For total protein 1 = 0.02, for albumin 1 = 0.0323 (i.e. $\frac{1}{3}1$)

Filmingen is calculated as follows

Pending of Standard > 0 15 × 2 > 100 × 625 Reading of Unknown

= grams fibringen per 100 ml of plasma

Globulin = Total Protein - (Albumin + Fibringen) in the case of plasma for serum Globulen = Total Protein - Albumin

For photometric measurement Calculations are the same except that the expression Density of Standard replaces the expression Reading of Standard used in the colormetine calculations. The limits of accurate colonmetric or photometric measurement are the same as those specified for the determination of nonprotein nitrogen on p-547 If an analysis falls outside of these limits, the digistion is repeated using a smaller or larger abquot as the ease may be, and the calculations are corrected accordingly

Interpretation. Normal values for plasma proteins, in g per 100 ml of plasma, are as follows albumin 46-67 globulin, 12 2.3, fibrinogen 0 3-0 6 A major function of the plasma proteins (see Chapter 22) is to aid in the normal distribution of water between the blood and tissues, albumin being approximately twice as effective as globulin in this respect gram for gram Edema almost invariably occurs when the total plasma proteins fall below the critical level of 5 3 g per 100 ml Increased plasma protein levels are noted in dehydration, due to diminished fluid intake or pathologic loss of fluid from the body (diarrhea, vomiting, surgical or traumatic shock, excessive burns Addison's disease), or when there is an absolute increase in globulin content, as in various anaphylactic conditions, malignancy, hver cirrhosis, and certain chronic infections Decreased plasma protein is found after loss of plasma by extravasation or renal excretion (albuminuria) or when protein synthesis is impaired owing to malnutation, vitamin deficiencies or diseases involving the digestive organs or liver Fibrinogen values are increased in pneumonia and other infections accompanied by kukocyto-is or suppuration, but are low in acute yellow atrophy of the liver poisoning due to chloroform or phosphorus and typhoid fever High sedimentation rate of red blood cells 15 associated with increased fibringen values in plasma

2 Method of Kingsley. "Principle The diluted plasma or serum is treated with a special buriet reagent. The color developed is compared with that of a standard protein solution treated similarly. By suitable fractionation of serum before treatment with the buriet reagent, albumin may be determined separately. From the total serum protein and albumin contents, globulin is determined by difference. The method as described has been slightly modified to permit the determination of total protein, albumin, and globulin on the same sample of serum.

Procedure 18 Transfer 0.5 ml. of unhemoly zed serum to a graduated centrifuge tube, or smail (13 × 125 mm.) test tube graduated at 10 ml. Add 23 per cent sodium suifate solution to the 10-ml. mark. Mix well by repeated inversion (but do not shake), and immediately pipet out a 2-ml portion of the uniform suspension and place in a separate small test tube, which need not be graduated This is Tube 1, which will be used for the determination of total protein To the remainder of the suspension in the centrifuge tube add about 3 ml of ether, stopper, and shake vigorously. Centrifuge for about 5 minutes, capping the tube to prevent loss of the ether. After centrifuging, hold the tube in a slanting position and insert the tip of a 2-ml transfer pipet of narrow bore along the side of the tube past the white layer of packed globulin precipitate and into the clear fluid below. Remove a 2-ml portion of the clear fluid with the pipet, wipe off any precipitate and entries to the outside of the pipet, and transfer to a second test tube (Tube). This will be used for the

¹⁰ Kingder J Lab Clin Ved., 27, 840 (1942) See also Robiason and Hogden J Biol Chem 135 707 727 (1940) Wield ibid 157, 173 (1945) An improved hurrer reagent and procedura has been described by Weichselbaum Am J Clin Pathol 7, 40 (1946).

114 Reagents Required 23 Per Cent Sedium Sulfate Solution Dissolva 230 g of anhydrous reagent grade sodium sulfate m 600 to 700 ml of water in n beaker by heeting and stirring While still warm transfer to a 1 liter volumetine flash dilute with water to the mark, and mix Transfer to a clean bottle and place in the incubator or water hath at 37 C Keep at this temperature at all times stoppered to prevent evaporation since some of the salt will crystallize out if keet at room temperature.

Special Biurd Reagant Prepare a saturated solution of sodium hydroxids which is can bonate-free (see Appendix This saturated solution should contein about 75 g of sodium hydroxide per 100 ml after praparation. This may be checked by titration of a diluted por ton Measure 92 ml of the carbonate-free saturated solution of sodium hydroxide containing 69 g of sodium hydroxide into a 500-ml graduated cylinder and add water to the 300-ml mark. Add 100 ml of a 1 per cent solution of crystalline copper sulfate and stit to mix Transfer to a clean hottle fitted with a rubber stopper. This reagent is stable for months at committee the solution of the stable contains of the stable contains and the stable contains a stable for months at the stable solution of the stable solution of the stable contains the stable for months at many stable solutions.

Standard Protein Solution Obtain a pooled for of normal human serum (10.15 ml will be needed) Determine the total protein content of a portion by micro Lyeldahl or mecro-Kieldahl analysis (see p. 001). Ditute 5 ml of the remainder to 100 ml in a volumetric flask with 15 per cent sodium chloride solution and mir Label with the total protein content in grains per 100 ml of the original serum since this value is used in the calculations. This solution is usable for about a month if kept in the refrigerator at all times. A new solution may be standardized by colorimetric comparison with the previous standard if deterioration of the latter has not occurred. If this procedure is followed check occasionally by the Kipt daily included to avoid the possibility of error. Since with a particular photometer the density of a given standard is quite constant if the analysis is properly cerried out this is as good an index as any of possible changes in the standard.

For colorimetric measurement three colored standards are prepared representing the burst reaction on seria at level of approximately 3 6 and 0 g per cent total protein resectively. These standards are stable for at least one month if stored in the efficient of blandard I To 1 ml of serium of kinwan protein content add 30 ml of water and 80 ml of the burst reagent Standard II To 3 ml of serium add 37 ml of water and 80 ml of burst reagent 1 standard II To 3 ml of serium add 37 ml of water and 80 ml of burst reagent 1 me equivalent protein values for these standards used in the calculation are as follows Standard I C/2 Standard II C Standard II 3 C/2, where C is the total protein content of the serium used in grains per 100 ml

albumin determination. For photometric measurement, place 2 ml of the standard protein solution in a third tube, and 2 ml of water alone in a fourth tube as a blank. For colorimetric measurement these latter two tubes are unnecessary since the unknown is compared against standards which have already been prepared (see below and footnote 165). To each tube add 4 ml of the special bimet reagent, followed by 2 to 3 ml of ether Stopper, shake vigorously, and centrifuge for 5 minutes. By means of a pipet, transfer the solution under the ether layer to colorimeter cups or photometer cuvettes and read within the next 10 to 20 minutes, preferably as soon as possible since on prolonged standing a slight turbidity may develop which requires further treatment with ether and centrifugation. For photometric measure ment, set the photometer to zero density with the blank, at 520 mm

If the total protein content alone is desired, the fractionation with addium suifate solution is omitted Dilute 05 ml of acrum to 10 ml with 09 per cent sodium chiloride solution, mix, transfer a 2 ml portion to a small test tube, and continue as described for Tube 1 above Alternatively, 01 ml of serum, measured in a pipet calibrated "to contain," may be pipetted into 19 ml of 09 per cent sodium chiloride solution in a test tube, the pipet being rinsed several times with the diluting fluid, and this 2 ml portion treated with buret reacent as described above

CALCULATION For colorimetric measurement use a green filter (with maximal transmission at 520 ma) over the experce set the unknown at 10 mm and read sgamet those two of the three standards desembed in footnate 165 which appear on inspection to be lower and higher respectively than the unknown. Calculate results as follows

$$\frac{A \times \frac{P_L}{10} + B \times \frac{R_H}{10}}{2} = \text{grams protein per 100 ml}$$

where A and B represent the equivalent protein concentrations of lower and higher standards respectively in grams per 100 ml of serum and R_L and P_B are the respective standard readings If the unknown reading is within 10 or 15 per cent of either standard it e comparison with the other standard may be omitted in which case the calculation becomes

Reading of Standard
$$\times C \times f = \text{grams protein per 100 ml}$$
 of serum

where C is the total protein concentration of the standard serum used in grams per cent and f is 0.5 1.0 or 1.5 depending upon whether Standard 1.11 or III is used. Calculations for Tubes 1 and 2 are the same to value for Tube 1 surpain total

Calculations for Tubes I and 2 are the same the value for Tube I gring lotal protein in grams per cent and tle value for Tube 2 giving sibumin in grams per cent Globuline Total protein Albumin

For photometric measurement

Total Protein

1lbumin

$$\frac{\text{Density of Tube 2}}{\text{Density of Standard}} \times C = \text{grams slbumin per 100 ml serum}$$

[&]quot;If the photometer is equipped for use with small test tubes which can be centrifuged these n as be used for color development and subsequent reading without the necessity for transfer.

Chap 23

Total protein - albumin = grams globulin per 100 ml serum

C is the total protein content of the (undiluted) standard serum, in grams per cent 44.520 m μ and in a 1-cm cuvette a serum containing 7 g per cent of total protein has a density of approximately 0.400 when corrected for the density of the blank (Fig 157). Up to 11 g per cent of total protein may be determined accurately under the conditions described. For higher values or with deeper cuvettes use 1 ml portions instead of 2 ml portions in Tubes 1 and 2 plus 1 ml of water, and multiply the results by 2

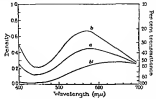


FIG 157 ABSORPTION SPECTRA OF COLORED SOLUTIONS OBTAINED IN KINGSLEY METHOD FOR PROTEIN

For hurst reagent alone (bl) and for hurst reagent in presence of serum containing 3.5 g per cent total protein (a) and 7 g per cent total protein (b) Solution depth 1 cm

Interpretation. Values for most normal and pathological sera by this method show good agreement with micro-Kipeldahl determinations. Discrepancies have been noted under certain conditions, but it is not known whether this is due to alteration in introgen content or in burier reacting power. The method is known to give satisfactory clinical results. For further interpretation, see the preceding neithed.

3 Determination of Plasma or Serum Protein, Hemoglabin, and Cell Volume (Hematocrit) by the Copper Sulfate-Specific Graity Method in Principle. The specific gravity of whole blood, plasma, or serum is established by allowing small drops of the material in fall into a series of copper sulfate solutions of known and varying specific gravity and noting whether the drops rice, fall or remain suspended under the defined conditions. From the specific gravity thus established, the plasma or serum protein, hemoglobin, and hematocrit values are obtained on the basis of an experimentally established relationship between these various quantities. If the plasma or serum protein content alone is in be determined, only plasma or serum is needed. Approximate hemoglobin and hematocrit values are obtained with a few drops of whole blood alone. Accurate hemoglobin and hematocrit values are obtained by determining the specific gravity of both whole blood and its plasma.

¹⁸⁷ As developed by Phillips V an Sly ke Dole Pmerson Hamilton and trelubrid at the Hospital of the Rockefeller Institute for Wederal Research 1 or a gradient tube method base I on a similar principle but using nonsqueous solvents in a single tube and suit-tle for years amaliamounts of serum see Lowyr and Hunter J Biol Chem 159 465 (1945)

Procedure. Venous blood is collected with a minimal amount of stasis (tourniquet applied for not over one minute). Capillary (finger tip) blood may be used, except in shock, for the approximate hemoglobin and hematocrit determination. To obtain plasma, transfer the blood immediately after drawing to a container having either a dried film of heparin sufficient to provide approximately 0.2 mg. for each mi. of hlood expected, or a dried film of the Heller and Paul oxalate mixture!** auffielent to provide not over I mg. per ml. of blood expected. Mix to dissolve, and centrifuge. If both whole blood and plasma are wanted, save a portion of the well-mixed blood for the whole-blood determination and centrifuge the remainder to obtain the plasma. For serum protein determination, serum from ordinary clotted blood is used.

Prepare a series of standard copper sulfate solutions of known and vary ing specific gravity.11 For the most accurate work, the series should be graded at intervals of 0 001 in specific gravity; for rougher work, intervals of 0 001 suffice. For plasma and serum, the range of specific gravity covered should be 1.015 to i 035; for whole blood, 1.035 to 1.075.

Allow a small drop of the sample to fall by gravity from a height of about i cm. above the solution (see Fig. 158) into one of the standard copper sulfate solutions having approximately the specific gravity



158 METHOD OF Fig DROPPING BLOOD INTO

expected. The smaller the drop the better; use a medicine dropper with a drawn-out tip, or a syringe needle, if the sample is whole oxalated or heparinized blood, the cells and plasma must be thoroughly mixed by repeated inversion immediately before use, otherwise serious error will result. Observe the behavior of the drop within the 10second period after it has lost the momentum of its fall. If the drop rises at all during this period, it is lighter than the test solution; if it continues to fall, it is heavier; if it remains stationary after momentum is lost, it has the same specific gravity as the solution After the 10-second period indicated, the behavior of the drop has no signifi cance, because of changes in specific gravity due to diffusion through the copper proteinate film around the drop. If the specific gravity of the drop is not established exactly by the first test, repeat the procedure with a fresh drop on solutions of higher or lower specific gravity as the case may be, until a solution is found in which the drop either remains stationary, which gives the specific grafity of the sample, or there are two adjacent stand-

COPPER SULFATE ards in one of which the drop rises, and in the other it falls in this case Interpolation between the values of the two standards is used; by noting the relative rate of rise and fall in the two solutions, it should be possible to interpolate to one-quarter of the difference in specific gravities between them Thus by experience one should be able to tell whether the specific gravity of the drop is halfway between the two standards, or nearer to one than to the other

Dissolve 3 g of ammonium oxalate and 2 g of potassium oxalate in water and dilute to 250 ml Pipet 0.55 ml of this solution for each ml of blood to be received in a container (small text tube or bottle) agreed in a film and dry in an incubator at 37° C or in a yacuum desiccator See Heller and Paul J Lab Clin Med 19, 777 (1934)

If oxalated whole blood and plasma are used, correct the observed values for the oxalate by subtracting 0 0004 for each mg of oxalate mixture present per ml of blood, applying the correction to both whole blood and plasma results Heparinized blood or serum requires no correction. Use the corrected values in obtaining results.

CALCULATION From the determined specific gravities, results are obtained by reference to the line chart shown in Fig. 159 For approximate hemoglobin and hematocrit

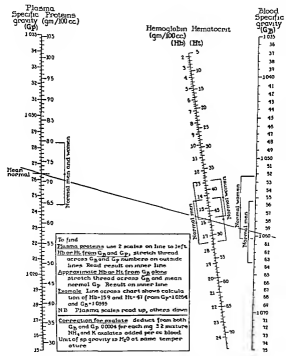


Fig. 159. Time Chart for Calculating Plasma Proteins, Hemoglobin, and Hamatocrit from Specific Grantiffs of Plasma, and Blood

determination using wi ole blood alone the blood is assumed to have a normal plasma specific gravity of 1 0264, and a line from this point on the left liand scale of the chart is drawn across the chart to the determined whole blood specific gravity as measured on the right-hand scale. The htmos, John content and hematocrit value are read from the interception of this line with the middle scale. Where both plasma and whole blood specific gravities have been determined the line is drawn connecting the two measured values on their respective scales and the bemoglohin and hematocrit values read from the intercept on the middle scale. For plasma protein content, in grams per 100 ml, the left hand scale is used If sorum is used instead of plasma the same scale is used the results then being in terms of grams of protein per 100 ml of serum

The copper sulfate solutions are permanent if kept stoppered to prevent evaporation. A given standard may be used over and over again, until one fortieth its vol ume of blood, plasma, or serum has been added 1e, one small drop per ml Thus a 100-ml portion serves for about 100 tests For routine work it is good practice to keep a record of the number of drops added to a given standard, and to discard st when the indicated limit las been reached. The solutions are self-cleaning precipi tated material ordinarily settling to the bottom. If any material should stick in the surface film either during or after an analysis it should be dislodged by tapping o removed with a wooden applicator stick before further use. The hemoglobin of whole bloo I will impart a greenish color to used standards, this does not impair their effec tiveness Temperature control is necessary only in the preparation of the standards as described in the Appen lix once the standards are prepared, further temperature control is ordinarily unnecessary

Interpretation. See previous methods

Other Methods The micro-Kieldahl method may be made more accurate by distillation followed by nesslerization or titration, gasometric determination of the ammonia may also be used See discussion of non protein nitrogen determination on p 548 The colorimetric method of Greenberg170 19 described in the eleventh edition of this book The determination of the specific gravity is the hasis of the 'falling drop method of Barbour and Hamilton, 171 which has found considerable clinical application For a turbidimetric method see Looney and Walsh

DETERMINATION OF HEMOGLOBIN

Introduction. In spite of the great chineal importance of hemoglobin determinations as a rule they are the most poorly conducted of all blood chemical analyses The majority of so-called office instruments in the hands of the average worker are known to give errors as high as 20 per cent The very fact that a method is adapted for small amounts of blood should presuppose accurate calibration and use of pipets, careful dilution and color match and frequent checking of both instrument and techni cian on blood of known hemoglobin content

The standard method for the determination of hemoglobin, and the one upon which almost all others are ultimately based is the determina tion by gasometric methods of the oxygen-binding power (oxygen capacity) of the blood. This method is described in detail in Chapter 21. An advan

tage of this method is that when properly carried out the results represent the functional hemoglohim of the blood, or that portion which is capable of carrying oxygen, ¹⁷² and do not include such nonfunctional pigment as methemoglobin, carhon monoxide hemoglobin, etc., as is the case with most of the other common methods Results by the oxygen capacity method may be expressed directly in terms of oxygen, as volumes per cent, i.e., ml. of hound oxygen per 100 ml. of blood, or they may be converted to grams of hemoglobin by making use of the empirically established fact that 1 g of hemoglobin is capable of combining with 1 36 ml. of oxygen under optimal conditions. ¹⁷⁴ Thus an oxygen capacity of 20 volumes per cent corresponds to a hemoglobin content of 20/1.36, or 14.7 g per 100 ml.

Another method for the determination of hemoglohin which is also suitable for standardization purposes, particularly in laboratories where gasometric equipment is not available, is to determine the total iron content of the blood. The iron content of hemoglobin has been accurately established at 0 340 per cent, ¹⁷¹ and the hemoglobin iron ordinarily represents 98 per cent or more of the total blood iron. Thus if the iron content is determined, the hemoglobin content may he accurately established. A simple and reliable colorimetric procedure for this purpose, developed by Wong, is described elsewhere in this section.

Direct colorimetric procedures for hemoglobin determination range from the most simple to the most complex, with a corresponding range in accuracy. They utilize either the color of the blood itself (duluted or undiluted as the case may be) or the color produced by treating blood with various reagents. The color is compared with, or measured in terms of, a standard color obtained from or representing a blood of known hemo-

a standard color obtained from or a globin content. These procedures are best used in conjunction with the ordinary laboratory colorimeter or photometer as described helow. For clinical purposes, a number of special "hemoglohinometers" of various types have heen designed. The Dare hemoglohinometer, one of the simplest of the clinical instruments, is based on companson of a film of undiluted blood of uniform depth with a red glass plate of graduated

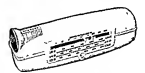


FIG 160 AO SPENCER HB-METER, Courtes), American Optical Company, Buffalo N Y.

intensity which has been snitably standardized. In the Spencer "Hb-Meter" (Fig. 160), the absorption of light by the hemoglobin in a layer of hemolyzed blood of fixed depth is compared with light absorption of a standardized glass wedge. The widely used Sahli type of clinical hemoglobinometer (Fig. 161), is based on treatment of the blood with dilute acid to produce the brown "acid hematin" color, followed by dilution until the color exactly matches a standard brown glass plate. From the

¹⁹ The desirability of this distinction may be limited under certain conditions, as for example in methemoglobinemia See also Rumington: Ann. Rev. Biochem., 12, 430 (1943).
¹⁹ By riphart and Skeges J. Biol., Chem., 147, 19 (1943).

dilution required, the hemoglobin content is readily obtained. This instrument is frequently very carelessly used, with care, however, it is capable of giving results accurate to within 5 per cent or so. The development of the photoelectric cell has produced hemoglobinometers of which the Fisher "Electro-hemometer" is an example (Fig. 162). This is contailly a one-purpose desk-model photoelectric photometer, the blood



Fig 161 Sauli Hemoolobinometer Courtesy Klett Manufacturing Co



FIO 162 FISHER LLECTRO-HEMOMETER. Courtesy Fisher Sc entific Co

sample, appropriately diluted in the glass tube is placed in the instrument and the hemoglobin content read directly from the scale, which is call brated by the manufacturers

For all hemoglobinometers supplied with a ready-made calibration regardless of type or source, the desirability of checking the calibration at intervals with blood of known hemoglobin content cannot be overemphasized. Manufacturers do not use these instruments, they merely make them, and the conditions of calibration and use may vary from time trine and differ from laboratory to laboratory. Only by careful checking under the conditions of actual use is it possible to climinate errors from this source.

It has been the custom among clinicians to express hemoglobin values in terms of per cent of some arbitrarily established normal. Unfortunately, there is no general agreement as to what the "normal" value should be for the very good reason that the hemoglobin content of the blood of a normal individual depends upon such factors as age, sex, occupation, climate, altitude and other environmental circumstances etc., and is obviously also influenced by the red cell count of the blood and factors which cause normal variation in this respect. The confusion attendant to the use of "per cut of normal" is illustrated by the fact that a blood hemoglobin content of say 145 g. per 100 ml., which is an exact value

10 ml of dilute hydrochloric acid Rinse the pipet thoroughly by sucking up the acid and blowing back several times. Allow to stand at room temper attree for at least 1 hour to complete color development, σ₁, if speed is essential, place in hot tap water (50 to 55° C) for 10 minutes ¹¹ Read in the colorimeter or photometer against the standard 'acid hematin 'solution. For photometric measurement, set the photometer to zero density with water or dilute acid as a blank, at 520 mμ

CALCULATION For colorimetric measurement

Reading of Standard Reading of Unknown $\times 0.075 \times \frac{100}{0.08} \times \frac{10}{100}$ = grams hemoglobin per 100 ml blood

In this calculation the 0 075 represents the hemoglol in content of the standard in 5per 100 ml the 0 05 is the volume of blood taken and the 10 (more exactly 10 05) represents the volume to which the blood is diluted If smaller or larger amounts of



FIG 163 ABSORPTION SPECTRA OF COLORED SOLUTIONS OBTAINED IN ACID HEMATIN METHOD FOR HEMOGLOBIN

For (a) 7 s g, per cent hemoglobin and (b) 15 g per cent hemoglobin each at a 1 200 dilution Solution depth 1 cm

blood are used at the same or different ddution volumes change the calculation accordingly. A blue glass or gelatin filter over the exepiece of the colorimeter improves the precision of color match.

For photometric measurement

 $\frac{\text{Density of Unknown}}{\text{Density of Standard}} \times 0.075 \times \frac{100}{0.05} \times \frac{10}{100} = \text{grams hemoglobin per } 100 \text{ ml blood}$

The acid hematin color which is really a colloidal suspension of hemin (see Chapter 22) has no characteristic peak light absorption in the visible spectrum (Fig. 163)

The choice of filter depends largely upon the sensitivity desired. At 520 m μ , and in a 1-cm. cuvette, the standard (equivalent to 15 g, per cent hemoglobin at a dilution of 1:200) has a density of approximately 0 500, and agreement with Beer's law is excellent over the entire range of hemoglobin content apt to be encountered.

This method has the advantage of a relatively permanent standard, in terms of which each unknown may be measured if will probably be found that for a given photometer and filter or wavelength setting the density of the standard will be constant and reproducible from day to day if mechanical or other changes do not occur in the photometer and if measurements are made under uniform conditions. In this event, the established density of the standard may be used in the photometer calculations without the necessity of reading the standard at each analysis. To eliminate error due to any changes in calibration, however, it is advisable to check the standard reading at intervals.

Interpretation. The hemoglobin content as determined by either the "acid hematin" or "alkali hematin" methods represents total hemoglobin, i.e., the methods do not distinguish hetween hemoglobin itself and such nonfunctional derivatives as methemoglobin, carbon monoxide hemoglobin, etc. Thus the results are apt to he slightly high as compared to oxygen capacity measurements, particularly in the ease of city dwellers and chronic tobacco smokers, whose blood may contain up to 5 per cent of its total hemoglobin as carbon monoxide hemoglobin. The "acid hematin" method as described is not applicable to the blood of species possessing nucleated crythrocytes (ie, hirds) because of the turndity caused by the nuclei; 100 the "akidali hematin" method does not have this disadvautage.

In normal individuals, at hirth the blood hemoglobin content may be over 21 g. per 100 ml. of blood, dropping to about half this value, or around 11 to 12 g per ceut, during the first six months after birth, and remaining at approximately this level up to about three years of age, after which there is a gradual increase until the age of 16 or so is reached. At this point, the hemoglobin content reaches a level which tends to he maintained throughout life; for males this level lies between roughly 14 5 and 16 0 g. per cent; for females, 13.5 and 15 0 g. per cent. As indicated on p. 613, the sex difference is due largely to differences in red cell count. This factor does not appear to explain the effect of age on hemoglobin content.

Aside from age and sex, other factors which influence the normal hemoglobin content include climate, altitude, evercise, menstruation, diurnal variation, and in general any factor which influences the red cell count or plasma volume A decreased hemoglobin content is found in the various anemias, in pregnancy, after moderately severe or chronic hemorrhage, and following the evcessive intake of fluids. An increased hemoglobin content, usually accompanied by an increase in red cell count (polycythemia) is found in hemoconcentration due to either shock or dehydration, in anoxia caused by either low ovgen pressure (high altitudes) or failure to properly ovgenate blood because of cardiac or pulmonary involvement, and evperimentally after the administration of cobnit safts.

¹⁵⁶ kor adaptation of the "acid hematin" method to chicken blood, see Schultze and Elvehjem: J. Biol. Chem., 105, 253 (1934).

- 2 Method of Neucomer 111 Principle The blood is treated with dilute hydrochloric acid to produce the brown acid hematin color described for the previous procedure This is then matched in a colorimeter against a brown class plate which has been spectrophotometrically standardized to correspond to the color obtained from a known amount of hemoglol in This method is relatively little used, having been replaced by the various photometric procedures
 - 3 Direct Photometric Method Principle Blood is diluted in weakly alkaline solution The color intensity is then measured in a photometer at 540 mm, and the hemoglobin content estimated from the established reading of a blood of known hemoglobin content under similar conditions. In the procedure described here, very dilute ammonia solution is used for diluting the blood 0 1 per cent sodium carbonate solution has also been proposed for this purpose 122 If the diluted blood is treated before reading with a source of carbon monoxi le gas such as ordinary illuminating gas the procedure becomes the more exact carbon monoxide hemoglobin method of Palmer 112

Procedure Collect 0 02 ml of blood in an accurate micropipet calibrated to contain Wipe off excess blood from the outside of the pipet, and transfer the blood to 5 ml of dllute ammonium hydroxide solution114 in a test tube Rinse the pipet with the solution by filling and emptying several times Mix, transfer to a suitable container if necessary, and read in the photom eter at 540 mu Set the photometer to zero density with a blank of the diluti ammonia solution alone For the Paimer procedure, bubble illuminating gas through the solution (in the hood) for about 30 seconds before reading in the photometer at 540 mu

CALCULATION Density of Unknown × F = grams homoglobin per 100 inl hlood F is a factor established by running the above-described procedure on a blood (known hemoglobin content Obtain a sample of normal human blood and determin its hemogloi in content by either the gasometric method (Chapter 24) or if gasometr equipment is not available by the method of Wong described on p 617 Treat dupl eate or triplicate portions of this standard blood by the procedure described abov From the determined density and the hemoglobin content calculate F as follow

F = Hemoglobin content in g per cent Density

Once established for a given photometer and filter or wavelength setting the value of F will ord narily 1 e constant unless mechanical or other changes occur in the photometer Checking the calibration at intervals will climinate errors due to change in the value of F In general the calibration factor for one instrument and filter is not applicable to another instrument or filter even of the same make

At 540 m_{\mu} and in a 1-cm cuvette a blood with 150 g per cent hemoglobin has 3 density of approximately 0 500 when simply diluted as descriled after earbon monox ide treatment the density is approximately 0 550 For spectrophotometric data, see Fig. 110 p. 473. Beer's law is valid over the entire range of hemoglobin content apt to be encountered For measurement using deeper cuvettes measure the blood into 10 ml. of ammonts solution instead of 5 ml establishing F on the same basis

Interpretation In the absence of significant amounts of abnormal blood pigments both the simple dilution procedure and the Palmer

¹⁶¹ Newcomer J Biol Chem 37 465 (1919) 55 569 (1923)

¹² Sanfo 1 Sheard and Osterberg Am J Clin Path 3 400 (1933)

¹ Ditte 4 ml f concertrated ammonium hydroxide to 1 liter with water and mls. able indefinitely

method are known to give satisfactory results. The Palmer method is more accurate, since it is free from any error due to the presence of carbon monovide bemoglobin in the blood as drawn. The chief disadvantage of the Palmer method in the past has been the difficulties associated with the preparation and maintenance of a sintable standard, this is eliminated by photometric calibration. The illuminating gas must of course be, free from substances other than earbon monovide which are capable of reacting with hemoglobin. If the blood contains significant amounts of methemoglobin, results will be in error. For the photometric determination of methemoglobin and total hemoglobin in the presence of methemoglobin, see the method of Evelyn and Malloy (p. 619). For other aspects

4 Method of Wong ¹⁸⁵ Principle The iron is detached from the hemoglobin molecule by treatment with concentrated sulfurn acid in the presence of potassium persulfate, without heating After removal of the proteins by tungsta caid, the iron in the filtrate is determined colorimetrically. From the total iron content, the hemoglobin content is readily obtained, since hemoglobin contains 0.34 per cent of iron, and only about 1 to 2 per cent of ess of the total blood iron is nonhemoglobin more

of interpretation, see previous methods

Procedure ³⁸⁸ With on Ostwold or micropipet, accurately transfer 0.5 ml of iron-free concentroted sulfuric acid. Mix by whirling one to two minutes Add 2 ml of Iron-free concentroted sulfuric acid. Mix by whirling one to two minutes Add 2 ml of saturoted potossium persulfate solution. Mix and dilute to about 2.5 ml with water Add 2 ml of 10 per cent sodium tungstate solution. Mix Cool to room temperoture under the tap, and dilute to volume with water. Stopper ond mix by inversion. Filter through a dry poper, collecting the filtrate in a dry flash. Prepare o standard in o second 50 ml volumetric flash by adding to obout 25 ml of water in the flash the following 2 ml of concentrated sulfuric acid, 2 ml of saturated potassium per sulfate solution, and 2.5 ml of standard iron solution containing 0.1 mg

of ferric iron per mi Cool to room temperature, dllute with water to the mark, and mix For photometric measurement, prepare a blank similar to the standard except that the standard fron solution is omitted.

Measure 10 ml of unknown filtrate, standard, and blank if necessary, into separate test tubes To each add 0 5 ml. of saturated persulfate soin tion followed by 2 mi of 3 N potassium thiocyanate solution Mix hy inversion and read in the colorimeter or photometer within the next 30 minutes For photometric measurement, set the photometer to zero density with the blank, at 480 mu

An alternative procedure for color development using thiogiycoliic acid is as follows. To the 10 mi portions in small flasks or test tuhes as described above, add 0 i ml of thioglycollic acid, 167 and 2 ml of concentrated am monium hydroxide Nix well, cool to room temperature by placing in cold water, and read in the colorimeter or photometer at any time within the next hour Just before reading, shake again, the color may fade on prolonged standing but it can be immediately restored by shaking in air For photometric measurement, set the photometer to zero density with the hlank, at 540 mu

CALCULATION For colorimetric measurement, using either the thiocyanate or thio-Liveolic acid procedure

Reading of Standard $\times 0.25 \times \frac{100}{0} \times \frac{1}{3.4} = \text{grams hemoglobin per 100 ml hlood}$

The value 1/3 4 represents the fact that 1 g of homoglobin contains 3 4 mg of iron. If this factor is omitted in the calculations the result gives mg of total iron per 100 ml of blood Under ordinary circumstances, less than 2 per cent of the total blood iron is from sources other than hemoglobin, this nonhemoglobin iron is therefore neglected in the calculation of hemoglobin content or a suitable correction may be made

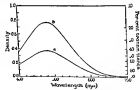


FIG 164 ABSORPTION SPECTRA OF COLORLO SOLUTIONS OBTAINED IN WONG METHOD FOR 1BON AND HEMOGLOBIN

For standards containing (a) 123 mg. and (b) 2 amg iron in 30 ml Solution depth 1 cm

For photometric measurement using either procedure

Density of Unknown Density of Standard $\times 0.25 \times \frac{100}{0.5} \times \frac{1}{3.4}$ = grams hemoglobin per 100 mL blood

The thiocyanate color has maximum absorption at about 480 m_µ (Fig. 164). At this

ist Burmester (J. Biol. Chem. 105, 189 (1934) adds sodium sulfite solution prior to add ng the thogher like acid to reduce the persulfate. This does not appear to be necessary

wavelength and in a 1-cm cuvette the density of the standard described is approximately 0.750. Since the standard corresponds to a blood iron content of 50 mg per cent equivalent to 14.7 g per cent bemoglobin all values of hemoglobin ordinarily encountered may be satisfactorily determined under these conditions. With deeper cuvettes use a a ml aliquot of filtrate instead in 10 ml add 5 ml of the reagent blank solution used to set the photometer to zero density develop and read the color as described and multiply the results by 2. If a filter at 450 mµ is not available equally satisfactory results at about 40 per cent lower scale reading may be obtained at 420 or 520 m.

The thioglycollic acid color has maximum absorption at 540 m μ . As compared to the thiocyanate color the density of the standard under similar conditions is considerably less being approximately 0.240 at peak absorption. This is not too low for accurate photometric measurement however and the color is superior in many respects to the thicoyanate color for analytical purposes. Excellent agreement with Beer's law is found up to double the standard concentration at almost any way elength between 400 to 580 m μ . The only disadvantage appears to be a tendency of the color to fado on standing as already mentioned restoration to the original value is easily accomplished by brief shahing in air.

Interpretation. This method is recommended for standardization of other hemoglobin procedures in the absence of facilities for determining oxygen capacity For other aspects of interpretation, see previous methods

5 Photometric Determination of Methemoglobin and Total Hemoglobin (Method of Evelyn and Malloy) *** Principle** \ \text{Nethemoglobin has a characteristic hight absorption at 635 mμ (see Fig 110 p. 473) this absorption is abolished in the presence of cyanide which converts methemoglobin to cyanomethemoglobin. The difference in light absorption at 635 mμ before and after adding cyanide is a measure of the methemoglobin present Total bemoglobin as determined by converting all the bemoglobin present to cyanomethemoglobin and measuring the light absorption at 340 mμ. Evelyn and Vialloy also desembe a procedure for measuring sufficemoglobin with the same sample this is omitted here because of the uncertainty associated with standardization For details see original article.

(a) METHEMOGLOBIN Determine the density of the solution in a photometer at 635 m_H, setting the photometer to zero density with water. This is reading D₁. Add 1 drop of neutralized sodium cyanide to the entire 10-ml, sample, mix, allow to stand 2 minutes, and make a second reading, D₁, under the same conditions as the first reading. The difference between D₁ and D₂ is the measure of the methemoglobin content which is calculated as described below. Any slight turbidity present is immaterial, since it is the same in both readings.

(b) TOTAL HEMOGLORIN. Use either the sample treated with cyanide as just described, or the original 1 101 dilution of the blood if total hemoglobin alone is to be determined. Add 1 drop of concentrated ammonium bydros ide to the entire 10-mil, sample to clear It, mix, and transfer a 2-mi, portion to a test tube containing 8 mil, of M/15 phosphate buffer at pil 6 6 and 1 drop of 20 per cent potassium ferric, ander. Mix and allow to stand 2 mustes to convert hemoglobin to methemoglobin. Add 1 drop of 10 per cent sodium cyanide, mix, and again allow to stand 2 minutes for the formation of cyanomethemoglobin Determine the density in the photometer at 540 mµ, setting the photometer to zero density with a blank consisting of 10 ml of M/5 phosphate buffer plus 1 drop each of the ferricyanide and the sodium cyanide solutions. Let this reading be D₁.

CALCULATION (a) Velhemoglobin

$$(D_1-D_2)\times F_M=$$
 grams methemoglobin per 100 ml. blood

where F_M is a factor expressing the relationship between a known amount of methomoglobin and the change in density at 635 m_H after adding cyanide. This factor is established as described below.

(b) Total Hemoglobin

$$D_4 \times F_T = {\rm grains}$$
total hemoglobin per 100 ml. blood

where F_{T} is the calibration factor for hemoglobin as cyanomethemoglobin, determined as described below

DETERMINATION OF CALIBRATION FACTORS ¹³⁹ Obtain a sample of normal human blood and determine its hemoglobin content by either the gasometric method (Chapter 24) or the norm method of Wong (p. 617). Of the well mixed blood, transfer 0 1 mt to a tube containing 9.9 ml of M/60 phosphate buffer at pH 6.6 and 0.1 ml of 5 per cent potassium ferrice andle solution. Mix allow to stand 2 minutes, and then defermine the density (= D₁) in the photometer at 635 mg, setting to zero density with blank consisting of 10 ml of the phosphate buffer plus 0.1 ml of the ferricande Viter the reading has been made add 1 drop of neutralized sodium eyands solution to both the entire sample, and blank mix and allow to stand 2 minutes and again determine the density (= D₂) against the blank at the same wavelength. The factor F_M is calculated as follows

The factor is therefore

$$\frac{152}{(0357 - 0055)} = \frac{152}{0302} = 504$$

To obtain F_T dilute a 2 ml portion of both the ferricyanide-eyamide treated stand and and blank to 10 ml with water and mix. Set the photometer to zero density at 540 m_{μ} with the blank and determine the density of the standard (= D_s)

$$F_T = \frac{\text{Hemoglohin content of standard in g per cent}}{D_1}$$

To illustrate the D_2 reading for the blood containing 15 2 g per cent hemoglobin was 0 211 Therefore $F_T = 15$ 2/0 211 = 72 0

Once they have been carefully established for a particular photometer and filter or wavelength setting these factors should be valid indefinitely unless mechanical or other changes occur in the photometer. Checking at intervals will eliminate errors due to such changes.

Interpretation The limit of precision of both the methemoglobin and total hemoglobin methods as described here is about 01-02 g per cent Using these methods with a Beckman spectrophotometer, in a study of 14 normal young men daily over a four day period it was not possible to find even 0.1 g per cent of methemoglobin in any except three random samples Paul and Kemp, 191 using a similar procedure but at a lower dilution of blood (and therefore possibly increased sensitivity), claim that small amounts of methemoglobin (0 03 to 0 13 g per cent) are regularly present in normal blood Increased methemoglobin content (methemoglobinemia) is associated with the administration of a variety of drugs, such as nitrites, aniline and derivatives sulfanilamide acetaunlide, etc When the drug is discontinued, the methemoglobinemia begins to decrease, it is thought that methemoglobin can be converted back to hemoglobin in the red cell, hence any evident methemoglobinemia presumably represents a balance between rate of production and rate of reconversion to hemoglobin 192

The cyanomethemoglobin method for total hemoglobin is considered to be one of the most accurate of the colorimetric methods. Results will be be one of the most accurate of the colorimetric methods. Results will be slightly in error in the few instances where the relatively rare pigment sulfhemoglobin is present, Evelyn and Malloy (loc cit) discuss this possibility of error and describe a correction for it. For further aspects of interpretation, see previous methods.

Other Methods. Hemoglobin may be quickly and accurately determined along with the total plasma protein, by the copper sulfatespecific gravity method (p. 607). The use of pyridine hemochromogen has also been advocated as a basis for colorimetric or photometric measurement. Manual the advantages claimed is that the method may be advantages elaimed in Ultramicromethods based on the benizidine color test for blood (see Chapter 22) have been described

¹⁹¹ Paul and hemp Proc Soc Expl Biol Med 56 55 (1944)

Cox and Wendel J Biol Chem 143 331 (1912)
 Rhimington Brit. Med J 1, 177 (1944) Phink and Watson J Biol Chem 146 171 (1944)

by Wu¹⁹⁴ and by Bing and Baker ¹⁹⁵ For the colorimetric determination of carbon monoxide hemoglohin, see p 195 Gasometric procedures for the determination of hemoglobin, carbon monovide hemoglobin, and methemoglobin are described in Chapter 21

DETERMINATION OF LACTIC ACID

Introduction. Blood lactic acid is ordinarily determined by conversion to acetaldchyde, which is then measured by titrimetric or color metric methods. A gasometric procedure based upon oxidation with permanganate to produce earbon dioxide has also been described 198 Of the various methods, the colorimetric method described here is by far th most sensitive, being readily applicable to 0.1 ml or less of blood 1 obtaining blood for lactic acid determination, precautions must be observed against the conversion of blood glucose to lactic acid on standing (gl_colysis) The use of fluoride as anticoagulant (see p 541) will prevent glycolysis, if oxalate or heparin are used, the protein free filtrate should be prepared as soon as possible after drawing the blood 197

1. Method of Barker and Summerson 112 Principle The glucose and other interfering material of the protein free blood filtrate is removed by the Van Slyke-Salkowski method of treatment with copper sulfate and calcium hydroxide An aliquot of the resulting solution is heated with concentrated sulfuric acid to convert lactic acid to acetaldehyde, which is then determined colorimetrically by reaction with p-hydroxydiphenyl in the presence of copper ions

Procedure 111 Deproteinize the blood sample (whole blood, plasma) with either tungetic acid (p 543), trichieroacetic acid (p 631), or zinc hydroxide

Wu J Biochemistry (Japan) 2 189 (1922)
 Bing and Baker J Biol Chem 92 589 (1931) See also McFarlane and Hamilton Brochem J 26 10-9 (1932)

Avery and Hastings J Biol Chem 94 273 (1931)
 Friedemann and Haugen (J Biol Chem 146 67 (1942)) describe a procedure in which
 Guid tl e drawn blood is ejected directly from the syringe into the protein precipitating fluid

¹⁸ Barker and Summerson J Biol Chem 438 535 (1941) ¹⁹ Reagents Required 20 Per Cent Copper Sulfate Dissolve 400 g of CuSO, 5H-O in about 1 liter of water with the and of heat cool, dilute to 2 liters and mix Stable indefinitely 4 Per Cent Copper Sulfate Dilute 1 volume of 20 per cent copper sulfate solution to 5 vol umes with water and min. Store in a bottle fitted with a stopper carrying a 1 ml pipet which lelivers approximately 20 drops per ml if this is done 1 drop may be used instead of the 0.05-ml portion specified in the text

Calcium Hydroxide Powder Either the USP or CP grades are satisfactory It is con vemently dispensed with a spoon spatula known to hold approximately 1 g. since exact measurement is unimportant

Sulfuric Acid Concentrated. Reagent grade iron free sulfuric acid is satisfactor. It is depensed from a buret suitally protected against absorption of atmospheric moisture The burst stopcock is cleaned the roughly of grease and lubricated with a little of the and theil in delivering pressures we roughly of grease and lubricated with a little of the ac-tiveli in delivering pressures should be taken against error due to the slow drainings the viscous and According to Russell (J. Biol. Chem. 154, 463 (1944)) intrates and mixing in the and will interfere only grades with how writers. in the acid will interfere only grades with low nitrate content are selected and if a par ticular lot shows poor color development it is discarded

p-Hydroxydiphenyl Reagent Dissolve 1 o g of p-hydroxydiphenyl (obtainable from Eastman Kodak Co. Richester N.Y.) in 10 ml of 5 per cent sodium hydroxide solution due a little water by warmers and arrange arrange and arrange arrange and arrange arrange and arrange arrange and arrange arrange and arrange arrange arrange and arrange arrange arrange and arrange arrang plus a little water by warrang and attring and dilute to 100 ml with water Store in a brown bottle fitted with a stopper and pipet capable of delivering 20 drops per ml. If this is done 2 drops may be used instead of the 0 1 ml portion specified in the text. The reagent is stable for many month's deterioration is evidenced by high blank readings

Standard Lacite Acid Sol dion. This is prepared preferably from lithium lactate which is

(p. 572) at a 1:10 dllutlon, Transfer 2 ml, of the protein-free filtrate, representing 0.2 ml, of blood, to a centrifuge tube graduated at 10 ml, in a second similar tube place 5 ml. of standard factic acid solution, containing 0.01 me, of lactic acid per ml, in a third tube place a little water; this is a blank. and serves to control the small amount of color yielded by the reagents alone. To each tube add 1 ml. of 20 per cent copper sulfate solution and dilute to the 10-ml, mark with water. Add 1 g. of powdered calcium hydroxide to each tube, stopper,200 and shake vigornusly until the solids are uniformly dispersed. Allow to stand for one-half hour, repeating the shaking at least once in the interim. Centrifuge down the precipitate, and transfer duplicate I-ml, portlons of the supernatant from each tube to thoroughly clean and dry test tubes having an internal diameter of 18 to 23 mm.201 To each tube add 0.05 ml. of 4 per cent copper sulfate solution, followed by 6 ml. of concentrated sulfuric acid from a buret. The sulfuric acid should be added drop by drop at first, mixing the contents of the tube well during the addition. The tube contents will become hot; it is not necessary to cool the tube, After the acid has been added to all the tubes, place them upright in boiling water for 5 minutes, then transfer the tubes to cold water (preferably running) and cool to 20° C, or below. When the contents of the tubes are sufficiently cool (but not before) add 0.1 mi, of the p-hydroxydiphenyl reagent, drop by drop, to each tube. The reagent precipitates out on entering the concentrated acid: It is dispersed throughout the solution as quickly and uniformly as possible by lateral shaking. When the reagent has been added, place the tubes in a beaker of water at 30° C, and allow to stand for 30 minutes or longer. Redisperse the precipitated reagent at least once during this period. Finally place the tubes in vigorously boiling water for exactly 90 seconds, remove, and cool in cold water to room temperature. Transfer the colored solutions to suitable containers and determine the photometric density at 560 mu, using water for setting the photometer at zero density.

CALULATION. Average the duplicate results on the blank to obtain the blank density. Subtract this value from the averages of standard and unknown to obtain their true densities. Since the 1-ml portion of copper-lime supernatant used for color development contains 0 005 mg of lactic acid in the case of the standard, and this represents 0 02 ml of original blood in the unknown (i.e., a dilution of 50), the calculation in this case is as follows.

Density of Unknown Density of Standard × 0 005 × 50 × 100 = mg lactic acid per 100 ml blood

anhydrous (For the method of preparing pure lithnum lactate see Chapter 31) For the stock standard, dissolve 0 213 g of pure dry lithnum lactate in about 100 ml of water in a 1-liter volumetric flask, add about I ml. of concentrated sulfure acid, dilute to the mark with water, and mix This solution contains I mg of lactic acid in 5 ml, and is stable indefinitely if kept in the refigerator To prepare the working standard, dilute 5 ml of stock staudard to 100 ml in a glass-stoppered volumetric flask with water and mix. This solution contains 0 01 mg of lactic acid per ml, and is best prepared fresh dady.

³⁹⁹ Glass-stoppered tubes may be used or, if these are unavailable, cover the mouth of the with a small square or "Parafilin," fresh surface down, held in place by the finger tip "Parafilin" may be obtained from laborators supply houses. In this and other phases of the procedure, contact of the solutions with the skin must be avoided because of the possibility of contamination with lactic acid from the akin surfaces.

³¹ Wide test tubes are specified to facilitate thorough mixing by literal shaking, this is more important than usual because of the viscosity of the concentrated and used as solvent. After use, the tubes are best cleaned by simply mixing in liet tap water, followed by discussed with different partial properties should be cleaned with eliromic and cleaning mixture. Hot soaps water followed by thorough runsing with distilled water is abequate.

It is recommended that the average blank density be determined separately as described to minimize error due to possible variation in the blank. If desired a single blank tube may be run and used for setting the photometer to zero density, in which case the average measured densities of standard and unknown are used directly in the calculation. The blank color is ordinarily about 10 per cent of the standard color In a 1 cm cuvettle at 560 m_p the standard described (equivalent to 25 mg per cent blood lactic acid) has a density of approximately 0 400 (I ig 165) and up to 60 ms per cent blood lactic acid may be accurately determined. For higher values, or with deeper cuvettes use smaller aliquots of filtrato and standard for the copper line treatment but keep the final volume at 10 ml at this stage using the same amounts of 20 per cent copper sulfate solution and calcium hydroxide as described. Correct the calculations as necessary

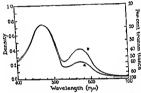


Fig 165 Absorption Spectra of Colored Solutions Obtained in Barker Summerson Lactic Acid Method

For standards containing (a) 0 002 mg and (b) 0 004 mg lactic acid Solution depth 1 cm

Interpretation. Venous blood of normal individuals in the resing state contains 5 to 20 mg of lactic acid per 100 ml ²⁰² During sever everuse this may rise to well over 100 mg per cent, decreasing rapidy during recovery Pathologically, increased blood lactate content is noted in general whenever there is a deficient supply of oxygen to the tissue (pneumona, beart failure) or the organism is unable to maintain the normal equilibrium between lactate production and utilization, as after the administration of anesthetics and in liver disease. Usually the presence of excessive amounts of blood lactate is at the expense of equivalent amounts of blood hearbonate, and an acidosis results. This disappears as the lactate is utilized

2 Method of Friedemann and Graeser 202 Principle The glucose of blood filtrates is removed with copper sulfate and calcium hydroxide the lactic and converted to acctaldehyde, which is then combined with sodium bisulfite. The bound sulfate is determined iodometrically, as in the method for urine (see p. 916)

Procedure. To 10 mi of Fulln-Wu or Somogyl filtrate representing 1 ml of blood add 2 ml of 10 per cent CuSO, and 2 ml of 5 per cent suspension of

To convert mg per cent lactic acid into milliequivalents per liter divide by 90 Biol Pireldemann and Graeser J Biol Chem 100 201 (1933) See also Wendel J Biol Chem. 101, 47 (1933)

Ca(OH): Shake at intervals for one half hour and centrifuge Of the filtrate 5 ml duplicates are taken for analysis as in the method for urine (page 916) using 0 002 N iodine For the blank treat 10 ml of 0 1 per cent glucose in the same manner as blood filtrate

Interpretation. See the previous method

Determination of Lactic Acid in Tissues, Tissue Extracts, Etc. Either of the procedures described above is suitable for the determination of the lactic acid content of various types of hological material For tissue analysis, precautions must be taken against post-mortem changes in lactic acid content, by prompt freezing in solid earhon dioude ("dry ice") or by adequate treatment with acid to destroy enzyme systems present Proteins present may be removed by any of the common methods For the colormetric procedure, an aliquot of the protein free finid containing 0.02 to 0.10 mg of lactic acid is treated by the copper-lime procedure described for blood For the iodometric procedure the aliquot sbould contain about 0.5 mg of lactic acid. This is diluted to 10 ml and treated as described for blood.

DETERMINATION OF CHLORIDES

Introduction. The chlorides of whole blood are distributed to the extent of about one-third of the total in the red cells and two-thirds in the plasma Plasma or serum is therefore ordinarily used for analysis, if whole blood were used, variations in red cell content would affect results out of proportion to their clinical significance. In obtaining plasma, excessive amounts of auticoagulants such as oxalate must be avoided. since they influence water and chloride distribution between cells and plasma, heparm does not have this effect. For precise work, where small changes in chloride content are of significance, blood should be collected under oil to minimize changes in carbon dioxide tension, since this also affects chloride distribution between cells and plasma (chloride shift, see "Role of the Red Cells," Chapter 24) This precaution is not necessary in routine clinical practice, but it is important that measurements be made carefully, preferably using volumetrie flasks for dilution as in the preparation of protein-free filtrates, because of the high chloride content of plasma or serum and the slight variations which are of significance

Chloride content is commonly expressed in terms of milligrams of sodium cbloride per 100 ml of sample It is more exact, and preferable, to express cbloride concentration in terms of milliequivalents of chloride per liter, since the major functions of chloride in the body are concerned with osmotic pressure regulation and acid-base balance. One milliequivalent of cbloride ion corresponds to 35 5 mg, or 58 5 mg of sodium chloride. A plasma with a chloride concentration of 585 mg per cent expressed as sodium chloride therefore contains 100 milliequivalents of chloride per liter. The general relationship between these two methods of expressing chloride content is as follows.

chioride content is as follow

1 Method of Whitehorn 264 Principle The chlorides are precipitated from th blood filtrate by means of silver nitrate in the presence of nitric acid, and the excess of silver titrated with standard thiocyanate solution, using ferric ammonium sulfate as a indicator

Procedure 2018 Pipet 10 ml of the Folin-Wu filtrate into a porcelain dish Add with a pipet 5 mi of the standard silver nitrate solution and stir thoroughly Add ahout 5 ml of concentrated nitric acid (sp gr 1 42), mlx, and let stand for 5 minutes, to permit the flocking out of the silver chloride. Then add with a spatula about 0 3 g of powdered ferric ammonium sulfate and titrate the excess of silver nitrate with the standard thiocyanate solution until the definite salmon red (not yellow) color of the ferric thiocyanate persists in spite of vigorous shaking for at least 15 seconds A microhuret should be used

CALCULATION Subtract the number of ml of theoryanate required from a 00 (= ml of silver mitrate added) and multiply the result by 100 to obtain the chloride content expressed as mg of sodium chloride per 100 ml. To express results in terms of milliequivalents of chloride per liter calculate as above and divide the result by a 80

Interpretation. Plasma or serum normally contains from 570 to 620 mg per 100 ml, expressed as sodium chloride, or 98 to 100 mili equivalents of chloride per liter The corresponding figures for whole blood are 450 to 500 mg per cent, or 77 to 86 milliequivalents per liter la creased plasma chloride is noted in nephritis, and this determination may aid in deciding whether or not salt should be restricted in the diet Decreased plasma chloride may occur in gastrointestinal disturbances associated with vomiting or diarrhea, in pheumonia, and in Addisoas disease Because of the close association between water and chloride marked disturbances in water distribution (hydration dehydration) may not necessarily be accompanied by changes from the normal plasma chloride level

2 Method of Scholes and Scholes 104 Principle The sample is intrated with standard mercuric nitrate solution at the proper acidity in the presence of diphen) carbazone as indicator Chlondes present react with the added mercuric ions to form soluble but undissociated mercuric chloride. When an excess of mercuric ion has been added, the indicator turns purple The end point is sharp and relatively stable

Procedure 101 Transfer 2 ml of Foisn Wu filtrate of plasma or serum (equif alent to 0 2 ml of original sample) to a small flask, and add 0 06 ml (4 drops)

²⁸⁴ Whitehorn J Biol Chem 45 449 (1921)

¹⁹ Reagents Required Standard Silver Virtale Solution Dissolve 2 905 g of cp silver nitrate in distilled water Transfer this solution to a liter volumetric flask and make up to the mark with distilled water. Mix thoroughly and preserve in a brown bottle. I ml. = 1 ms VaC! (It is to be noted that the silver nitrate and nitric acid are not added to the profess free filtrate simultaneously. To do so may result in the mechanical enclosure of silver mitrate solution within the curds and hence too high results)

Standard Thucyanate Solution Because these anates are lygroscopic the standard solution should be prepared volumetrically Dissolve about 1 7 g of KCNS or 1 4 g of VII.CNS in a liter of water. Titrate are 1 properly of the control of the co in a liter of water Titrate against standard silver nitrate solution under the conditions specified under I rocedure and dilute accurately so that 5 ml are equivalent to 5 ml of tl e silver mirate se lution

Solid ferrie alu n is used ratl er than a solution in order to insure a very high concentration in the mixture to be titrated. It is powdered to lacilitate its solution-144 Schales and Schales J Biol Chem 149 879 (1941)

²⁸⁷ Reagents Required Diphenylcarbazone Solution Dissolve 100 mg of s-diphenylcarba-

of diphenylcarbazone indicator solution. Titrate with the standard mercuric nitrate solution, using a microburet capable of being read to 0.01 ml. and delivering small drops. At the end point, the color of the solution changes from high vallow to deep number.

The plasma or serum may be titrated directly without previous deproteinization. This procedure eliminates errors in the preparation of the flitrate Transfer 0.2 ml of sample to a small flask, add 1.8 ml of water, 0.06 ml, of indicator, and titrate as above. The color of the solution undergoes several changes during the titration, becoming light yellow just before the end point is reached, and changing to pale violet at the end point. Results by the direct titration are slightly higher than when a filtrate is used, possibly because of slight loss of choude during deproteuration.

CALCULATION Results for either the protein free or direct titration are calculated as follows

ml mercuric nitrate solution used $\times \frac{100}{1}$ = milliequivalents chloride per liter

where 4 equals the number of ml. of mercuric nitrate solution required for 2 ml of standard sodium chloride solution. If 4 equals 200, the calculation simplifies to

mI mercuric nitrate solution used × 50 = milliequivalents of chloride per liter

If results are desired in terms of mg of sodium chloride per $100 \ \mathrm{ml}$ the calculation is as follows

ml $\,$ mercurie nitrate solution used $\times\,\frac{100}{A}\,\times\,5\,85\,$ = mg $\,$ NaCl per 100 ml

Interpretation. See the previous method This method, though not quite so accurate as the iodometric method described subsequently, is relatively simple and is known to give satisfactory results. It has definite advantages over the Whitehorn method. It may be applied to spinal fluid analyses, ordinarily without deproteinization. For the application to urine, see Adper, Schales, and Schales ¹⁶⁸.

3 Method of Sendroy, Modified by Van Slyke and Hiller. 109 Principle. The plasma or serum is treated with phosphone acid containing either tungstic acid or

zone (obtainable from Eastman Rodal, Co. Rochester, N.Y.) in 95 per cent alcohol and dulte to 100 ml. Store in a dark bottle on the cold. Loup the bottle with a rubber bulb medicine dropper whose tip is adjusted so as to deliver 65 to 70 drops of solution per ml Prepare fresh solution each month

Standard Sodium Chloride Solution Dry some reagent grade sodium chloride in an oven at 110° to 120° C overmith Cool and weigh out 534 5 mg Dissolve in a little water and transfer with runsings to a 1-liter volumetre flast. Dilute to the mark with water and nux This solution is stable indefinitely and contains 10 indhequivalents of chloride per little in 54 545 mg of standardize each new tot of standard.

mercuric nitrate solution

Stan lard Vercure A Wrate Solution Place a few hundred mi of water in a 1 liter volumet: the flask and add 20 mi of 2 N mire acai 4dd 3 g of reagent grade mercure intrate dissolve by shaking diduce to volume with water and mir Standardize as follows transfer 2 mi of standards sodium chioride solution to a small flask, add 4 drops of diphenylear ba zone solution and utrato with the mercurie intrate solution from a microburet as described in the text. The number of mi of mercurie intrate solution from a microburet as described in the calculations above If the strength of the mercurie nutrate solution is adjusted so that A equals 200 either by adding more increase interact or by dilution with water containing 30 ml of 2 N miru, send jer fiter as the case may be the calculations are simplified (see text) This standard mercurie intrates obtution is stable indefinitely and used not be protected from hight so that large amounts may be prepared at one time.

209 Sendroy J Biol Chem 120 335 405 419 (1937) ibid , 130, 605 (193J) ibid 142, 171

(1942) Yan 'lyke and Hiller (personal communication)

pieric acid, which precipitates the proteins. The mixture is then shaken with an excess of solid silver rodate and fiftered. Chlorides present react with the insoluble silver is rodate to form insoluble silver chloride, and soluble solate, which passes into filtrate. On the addition of solidate to the filtrate, the isolate reacts to produce free isolate, which is then titrated with standard thosulfate.

Procedure 33 Transfer 1 ml of plasma or serum to a 50 ml. Erlenmeyer flask Add 25 ml of phosphoric-tungatic acid out tion and mix To the mixture add 63 to 65 g of sliver lodate (measured with sufficient accuracy from a spoon apartula previously found to contain approximately the right amount). Stopper the flask and shake vigorously for 38 seconds. Pour onto a dry 9-cm. folded filter of loose texture, collecting the fittrate in a dry flask.

Phosphore Pure Acid Solution This may be used in place of the tungstic acid respect!

Chloride-free tungstate is not obtainable, pure and as less likely to contain chloride files

6 ml of phosphore acid and 2 g of pure and in a 1 liter flack add water to dissolve differ

to 1 liter and mix

Siler folds: This must be l'ea from mora soluble rodates. To test shake 0.5 gr. vith 25 ml of phosphorts tungatic acid solution for 1 minute filter and titrate 10 ml of of filtrate with 0.2308. NI tosulfate after doding rodate as described above for plasma filtrates. If more than 0.5 ml of throsulfate its required at a temperature of 20° cr 100 of the proper of the proper of the control of the proper of the control of the proper of the control of the proper of the control of the co

Sodium Iodide Solution Dissolve 50 g of reagent grade sodium rodule in 50 ml of water that is the solution must be desarded when a 1 ml portion added to 10 ml of phosphorto-tungsite acid solution must be dissarded when a 1 ml portion added to a 10 ml of phosphorto-tungsite acid solution gives a blue color in the presence of a diop of a starth solution For occasional analyses the dry reagent is a more economical A 0.5% portion is used for each analysis. Dispense from a apoon spatula previously found to deliver approximately the right amount.

Standard Sodrum Thosulfate Soliston Prepara a stock throsulfate solution by dissolving 57.3 g of crystaline sodium throsulfate VasSo3, 511,50 in water dutumn to 220 ml in 87.3 g of crystaline sodium throsulfate the Soliston of Soliston and Soliston Solist

¹¹⁸ Rosgents Required Phosphore-Tungstic Act I Solution Transfer 6 g of reagent grade chloride free sodium tungstate to a 1 liter solumetric flack add water to dissolve followed the segment-grade phosphore act of make up to 1 liter with water, and mix the state may be tested for cl lorde by dissolving 1 g in 10 ml of water, adding 20 m of contrated mitre acid and filtering into a test tube containing a fow in 0 if per cent sulve mitre if the filtrate does not show a precipitate of silver chloride the utility of the filtrate does not show a precipitate of silver chloride the utility of the filtrate does not show a precipitate of silver chloride the utility of the filtrate does not show a precipitate of silver chloride the utility of the contrated solution with an equal volume of 95 per cent alcohol filtering off the crystals from the cooled solution washing with alcohol and drying mixer.

Transfer 10 ml of the water-clear filtrate to a small flask Add I ml of sodium lodide solution (or 0.5 g of solid sodium rodide) Mix carefully with one or two swirls, and titrate the liberated iodine immediately with standard sodium thiosulfate solution. Add the thiosulfate in rapid drops, with careful mixing, until the yellow color of the iodine has almost disappeared. Add 2 drops of starch solution and continue titration until the blue color disappears and the solution is colorless, or, if picric acid was used for deproteinization, only the yellow color of the picric acid remains.

CALCULATION In this procedure, I equivalent of chloride ion leads to the production of 6 equivalents of iodine requiring 6 equivalents of thiosulfate for titration. Therefore I ml of N thosulfate represents 1/6 millieguivalent of chloride. Since the 10 ml of filtrate titrated represent 19/6 ml of sample the formula for calculation of results in terms of millieguivalents of chloride neg liter is as follows.

ml thiosulfate \times normality $\times \frac{1}{6} \times \frac{26}{6} \times 1000$

= millieguis alents chloride per liter

If the normality of the thiosulfate is exactly 002308, the calculation simplies to

ml thiosulfate × 10 = milhequivalents chloride per liter

If the thiosulfate is not exactly 0 02308 $^{\sim}$ and a factor is used as described under "Standardization," the calculation becomes

ml thiosulfate X factor X 10 = milliequivalents chloride per liter

To express results on the basis of milligrams of NaCl per 100 ml since 1 milli coursalent of chloride corresponds to 58.5 mg of NaCl the calculations become

ml 0 02308 N thiosulfate × 58 5 = mg NaCl per 100 ml

or

ml thiosulfate X factor X 58 5 = mg \nCl per 100 ml

Interpretation. See under the method of Whitehorn above This method may be used with Folin Wu filtrates (10 ml of filtrate made up to 26 ml with phosphoric-tungstic reagent, treated with rodate, filtered, and titrated exactly as described above, with the same calculations), it is directly applicable on an unchanged basis to the analysis of spinal fluid, gastric contents, urine, etc., and it may be used on a microscale with proportionate decrease in amounts of reagents used, and suitable change in calculations

4 Determination of Total Chlorides in Blood and Tissues (Von Slyke) 111 Principle. The proteins are oxidized, and the chloride is precipitated, by wet digestion with concentrated nitric acid in the presence of silver intrate. The excess silver is then titrated with thiocyanate, as described in Whitehorn's method

DETERMINATION OF PHOSPHORUS

Partition of Blood Phosphorus. Whole blood contains about 10 mg of total phosphorus per 100 ml, present clincily as unorganic phosphote, organic ocid-soluble phosphote esters, and lipide-phosphorus, with other phosphate-containing compounds possibly present in small amounts. The distribution of phosphorus between cells and plasma is quite uneven,

³¹¹ Van Slyke J Biol Chem., 58, 523 (1923) Wilson and Ball J Biol Chem., 79, 221 (1928)

for example, cells contain much more organic and total phosphate than plasma, while the morganic phosphate of whole blood is practically cutirely in the plasma For the determination of morganic phosphite and total acid-soluble phosphurus, prutern is precipitated with trichloroacetic acid and the filtrate used For lipide phosphorus determination, an alcohol-ether extract is obtained. In the determination of total acid-solu ble, lipide, and total phosphorus, organic matter is destroyed by digestion with sulfuric acid and 30 per cent hydrogen peroxide. The phosphate-contanning solutions thus obtained (as well as the trieliloroacetic acid filtrate used for the direct determination of morganic phosphate) are treated with molybdie acid, wherehy phosphomolybdie acid is formed from any inorganic phosphate present. On the addition of suitable reducing agents, phosphomolybdie acid is selectively reduced to yield a deep blue color ("molybdenum blue") which is apparently a mixture of lower oxides of molybdenum This color is used as a measure of the amount of pho-phate present

Bell and Doisy 212 earried out the reduction with hydroquurone in alkaline solution Briggs113 made the color more stable by using an acid solution Benedict and Theis'14 intensified the color by heating. This proceduro is satisfactory where only inorganic phosphate is present, as in serum or plasma filtrates, hut cannot be used on whole blood filtrates for example, since any phosphate esters present may be hydrolyzed by the heating in acid and thus lead to high values Fisko and Subballus 115 sug gested the use of 1,2,4 aminonaphtholsulfonic acid as reducing agent, at room temperature This method has been widely used, and is the one described here Kuttner and Cohen revived the original suggestion of Deniges that stannous chlorido be used as reducing agent, and this has hkewise found much favor Stannous chloride has the advantage over aminonaphtholsulfome acid that the stock reagent is quito stable and the color produced with phosphomolybdate is more intense, thus permitting the estimation of smaller amounts of phosphorus For example Shinowara, Jones, and Reinhart have described procedures for phosphate determination, using stannous chloride as reducing agent, which require as little as 0 06 ml of serum A disadvantage of stannous chloride is that the color intensity changes continuously with time, so that careful control of the time factor is essential, and deviations from Beer's law are found, usually requiring correction factors. In the authors' experience, the method of Fiske and SubbaRow has proved superior to any other method thus far described

1 Determination of Inarganic Phosphate (Method of Fiske and Subba-Rows Principle The proteins of blood are precipitated with trichloroacetic acid-The protein free filtrate is treated with an acid molybdate solution, which forms phos-

Bell and Dossy J Bsol Chem 44 45 (1920)
 Briggs J Bsol Chem 53 13 (1922) 59 255 (1924)
 Benedet and Thess J Bsol Chem 66 63 (1924)
 Fiske and bulballow J Bsol Chem 66 375 (1925)
 Fiske and Colen J Bsol Chem 75 317 (1927) also Kuttner and Lichtenstein J Bul Chem 86 671 (1930)

⁵¹² Sl mowara Joi es and ReinJart J Biol Chem 142 921 (1942)

phomolybdic acid from any phosphate present. The phosphomolybdic acid is reduced by the addition of 1.2 4-aminonaphtholsulfonic acid reagent, to produce a blue color whose intensity is proportional to the amount of phosphate present

Procedure 111 To 8 ml of 10 per cent trichloroacetic acid solution in a small flask, add slowly, with mixing, 2 ml of while blood, serum, or plasma Stopper, shake, and filter through a low ash filter paper Transfer 5 ml of filtrate to a cylinder or other container graduated at 10 ml, and add 1 ml of the Molybdate II reagent Mix Add 0 4 ml of aminonaphtholsulfonic acid reagent, and again mix Dilute to the mark, mix, and allow to stand 5 minutes

For colorimetric measurement, compare in the colorimeter against a standard prepared at the same time, as follows Transfer 5 ml of standard phosphate solution, containing 0.4 mg of phosphorus, to a 100 ml volumetric flask, and add from a graduate 50 ml of water Add 10 ml of Molyb date I (nor Volybdate II), mir, and add 4 ml of aminonaphtholsulfonic acid reagent Dilute with water to the 100 ml mark, mir, and allow to stand 5 minutes Compare the standard against Itself in the colorimeter before reading the unknown if the color of the unknown is particularly strong repeat the reading of the unknown a few minutes later, to be sure that maximal color development has taken place.

For photometric measurement, transfer a portion of the colored solution to a sultable container and read in the photometer at 660 to 720 m_µ (see Fig 166) Set the photometer to zero density with a blank prepared by treating 5 ml of 10 per cent trichloroacetic acid with 1 ml of Volybdate II and 0.4 ml of aminonaphtholsulfonic acid reagent, followed by water to a volume of 10 ml Establish the density of a standard phosphate solution as a follows Transfer 5 ml of the stock phosphate standard, containing 0.4 mg of P, to a 50 ml volumetric flask, make up to volume with 10 per cent tri chloroacetic acid, and mix Transfer 5 ml of this dilute standard, containing 0.04 mg of phosphorus, to a suitable container, add 1 ml of Volybdate If

² Reagents Required 10 Per Cent Trichlorocetic Acid Dissolve 10 g of reagent grade trichlorocetic acid in water and dilute to 100 ml
10 \(\nabla \) S \(\text{luric Acid Carefully add 450 ml of concentrated sulfuric acid to 1300 ml of

water To cheek dutte 10 ml of this solution to 100 ml in a volumetric flask mix and intrate a 10-ml portion with standard 1 N sodium hydroxide Prom the titration results adjust the original solution if necessary to make it exactly 10 N Mobilette 1. Dissolve 20 et al. research standard mixture in the property of the p

Molybdate I Dissolve °0 g of reagent-grade ammonium molybdate in about 200 ml of water in a 1 liter volumetric flask place 500 ml of 10 N sulfuric acid Add the molybdate solution and dilute with washings to 1 liter with water 'lix' Stable indefinitely

Molybdate II Dissolve 25 g of reagent-grade ammonium molybdate in about 200 inl of water ln a l liter volumetrie flask place 300 ml of 10 N sulfurie acid \dd the molybdate

solution and dilute with washings in I liter with water. Yux. Stable indefinitely. Aminonaphilohaldionic Acid Reagent Place 195 ml. of 15 per cent sodium bisulfite solution (see below) in a glass-stoppered cylinder Add 0.5 g. of 1.2 4-aminonaphilohaldionic acid (satisfactory material can be obtained from Eastman Kodsk Co. Rochester V. V. 4dd 5 ml. of 20 per cent sodium sulfite (see below). Stopper and shake until the powder is

hssolved If solution is not complete add more sodium sulfite 1 ml at a time, with slaking but a roid an excess Transfer the solution to a brawn glass bottle and store in the cold This solution is usable for about four weeks if kept as described to Per Cent Sodi im Britistle To 30 g of reagent-grade sodium bisulfite in a beaker add

²⁰⁰ ml of water from a graduated exhader Strr to dissolve and if turbil allow to stand cell-stoppered for several days and filter Keep well stoppere!

O Per Cent Sodium Sulfte Dissolve 20 g of reagent-grade anhi drous sodiu ii sulfite in water d lute to 100 ml and filter if necessary keep well-stoppered

So ulard Phosphate Solution D socket exactly 0.351 g of pure dry monopotassium photo hate in water and transfer quantitatinety to a 1 liter volumetric flash. Vid 10 in 10 f 10 \ sulfuric acid dilute to the mark with water and mix This solution contains 0.4 ing. of phosp to us in 5 in 11 to stall 10 in definitely

and 0.4 ml. of aminonaphthoisulfonic acid reagent, dilute to 10 ml. with water, and mix. Allow to stand 5 minutes and determine the density in the photometer whose zero is set with a blank as described above.

CALCULATION For colorinetric measurement Since the 5 ml of filtrate taken represent 1 ml of original sample, and the standard containing 0.4 mg of P is in ten times the volume of the unknown, the calculation becomes

 $\frac{\text{Re ding of Standard}}{\text{Reading of Unknown}} \times \frac{0.4}{10} \times 100$

- mg morganic P per 100 ml whole blood, plasma, or serum

For photometric measurement

Density of Unknown × 0 01 × 100

Density of Standard - mg inorganic P per 100 ml whole blood, plasma, or scrum

The density of the standard at 600 m μ in a 1-cm curvitie is approximately 0.500 (see Fig. 105), at 720 m μ , the peak position of the curve, the density is about 10 per cent

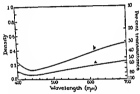


Fig. 166 Absorption Siectea of Colored Solutions Obtained in Fiske-SubbaRow Phosimate Method

For standards containing (a) 0.02 mg, and (b) 0.04 mg phosphorus Solution depth, 1 cm

greater Under these conditions the limit of accurate measurement corresponds to an inorgane phosphate content of approximately 7 to 8 mg per cent and this range is proportionately reduced with photometric measurement at greater depth of solution. To permit the covering of a greater range of phosphate concentration, the amount of sample taken for protein precipitation may be decreased e.g., 1 ml (or 0.5 ml) of whole blood scrum or plasma is treated with 10 per cent trichloroacetic acid at a final volume of 10 ml and 5 ml of filtrate taken for analysis as described II thus done the calculation is the same except that the final result is multiplied by 2 (or 4, if 0.5 ml of sample was taken). The color obtained in this procedure shows httle change between 5 and 20 minutes after adding the aminonsphitosluffor acid reager and the agreement with Beer's law is excellent permitting calculation of results in terms of the density of a simultaneously prepared standard and eliminating the necessity for a calibration curve

Interpretation. The normal morganic phosphorus content of the blood plasma or serum is about 5 mg per 100 ml in infants and children and 37 mg in the adult. In severe nephritis morganic phosphorus may

rise 15 to 20 mg, and may hear a relation to the acidosis found in such cases Clinically, the most important changes are those in rickets. In the rickets of children or in experimental rickets produced in animals by lowphosphorus diets the inorganie phosphorus of the blood may fall to 2 mg. or lower. Treatment with antirachitie vitamin or ultraviolet radiation increases the phosphorus of the blood and leads to recalcification of the rachitic lesions. Occasionally rickets may be accompanied by high inorganic phosphorus of the blood, and the tendency of irradiated ergosterol to restore normal phosphorus values may be independent of its calcifying activity. During the healing period following fracture of a hone, an increase in plasma phosphates is sometimes observed. The phosphate level of the plasma of children rises during the summer and falls during the winter. Rickets has its greatest incidence during the late winter, reaching its peak in March. These findings may be correlated with the degree of exposure to solar ultraviolet rays Injections of insulin decrease the phosphate of the plasma.

2. Determination of Total Acid-soluble Phosphorus: Principle. The organic matter in an aliquot of a trichloroacetic acid filtrate is destroyed by digestion with sulfutic acid and subsequent oxidation with 30 per cent bydrogen perconde The phosphate-contaming solution is then analyzed for phosphato by the method of Fiske and SubhaRow.

Procedure. 119 Prepare a trichlotoacetic acid filtrate of whole blood, plasma. or serum as described for the determination of Inorganic phosphate. Transfer 2 ml. of this filtrate to a test tuhe, 200 by 25 mm., and add 2.5 ml. of 5 N sulfuric acid and a quartz chip to minimize bumping. Piace in a slanting position over a microburner, with the burnet tip about 2 cm. below the bottom of the tube, or suspend in a wire hasket about 1.5 inches above an electric hot plate. After evaporation is complete and the mixture turns brown or hiack with no further change, remove the tube, cool slightly, and add 1 drop of 30 per cent hydrogen peroxide, allowing the drop to fail directly into the digestion mixture. Replace the tube and continue heating. The contents of the tube should become colorless; if not, repeat the addition of hydrogen peroxide and heating. When coloriess, cool the tube, add a few mi, of water, and heat to boiling momentarily. Coof again and transfer the contents of the tube to a 25-ml, volumetric flash, with washings until the flask is about half fuil. Add 2.5 mi. of 2.5 per cent ammonium molybdate solution, followed by i mi, of aminonaphtholsuifonic acid reagent. Dijute with water to the 25-mi. mark and mix. Allow to stand 5 minutes, then read in the colorimeter or photometer.

¹¹⁸ Reagents Required: 10 Per Cent Trichloroscette Acid Solution, Molybdate I Solution, Aminonaphtholsulfonic Acid Reagent, and Standard Phosphate Solution as described for the determination of Inorganic Phosphate (p. 631), and in addition.

⁵ N Sulfuric Acid, Till a 250-ml volumetric flask to the mark with 10 N sulfuric acid solution (see p. 631). Pour the contents of the flask into a 500-ml volumetric flask, with rinvings, and didute to the mark with water, Mix. Stable indefinitely.

^{2.5} Per Cent Ammonium Melybelate Solution Dissolve 2.5 g. of reagent-grade ammonium moly belate in water, transfer to a 100-ml, volumetire flask, fill to the mark, and mix. As soon as any appreciable amount of sediment forms in this solution, it should be discarded 50 Per Cent Hydrogen Peroxite, Only the highest purity, eventually phosphate-free marrial may be used. The products of Merck and of J.T. Baker are known to be satisfactory.

This reagent is extremely corrowing to the skin and must be handled carefully. Keep in the cold, and use a medicine dropper for dispensing

For colorimetric measurement, compare against a standard prepared as described above for the determination of blood inorganic phosphate. For photometric measurement, the most accurate procedure is to run a digested blank and a digested standard along with each series of unknowos This corrects for any phosphate or other factors in the reagents which may affect the final results Digested blank 2 ml. of 10 per cent trichloroacetic acid, treated with sulluric acid, evaporated, oxidized with hydrogen peroxide, and color reagents added, exactly as described for the unknown. Digested standard 1 ml of standard phosphate solution, containing 0 08 mg, of phosphorus, plus 2 ml of 10 per cent trichloroacetic acid, treated with sullunc acid, evaporated, oxidized with hydrogen peroxide, and color developed, exactly as described for the unknown. The color of the disested black and standard is developed at a final volume of 25 ml, as with the unknown Set the photometer to zero density with the black, and determine the densities of the standard and the unknown, using the same waveleogtb 25 specified for the determination of moreanic phosphate on p. 632.

CALCULATION For colorimetric measurement Since the standard contains 0.4 mE of P in a final volume of colored solution four times that of the unknown, and the 2 ml of unknown filtrate represent 0.4 ml of original sample, the calculation is as follows

Reading of Standard $\times \frac{0.4}{4} \times \frac{1}{0.4} \times 100 = \text{mg}$ total acid-soluble phot-phorus per likeding of Unknowo $\times \frac{0.4}{4} \times \frac{1}{0.4} \times 100 = \text{mg}$ total acid-soluble phot-phorus per 100 ml blood, plasma, or serum

If the reagents, particularly the trichloroacetic acid and the hydrogeo peroxide, contain significant amounts of phosphorus, the quantity present must be established by a blank digestion and analysis and suitable corrections made

For photometric measurement

Density of Unknown × 0.08 × 1/0.4 × 100 = mg total acid-soluble phosphorus per 100 ml blood: plasma, or serum

The conditions for photometric measurement have been presented in connection with the determination of morganic phosphate

Interpretation. The acid-soluble phosphorus content of normal blood is approximately 25 to 30 mg per 100 ml of whole blood. The total acid soluble phosphorus includes the morganic phosphate as well as those organic phosphate esters which are not present as hipide or nucleoprotein material. Such esters include hexosephosphate, diphosphoglyceric acid, "ecrtain free nucleotides, and other compounds which have not been characterized. In the plasma, the total acid-soluble phosphorus is represented largely by morganic phosphate. In the cells, the reverse is trepresented have a consideration of the relative volume of cells to plasma. There is some evidence that the acid-soluble phosphorus of hlood is of significance in acid-base balance in the body, since it is found to be considerably depleted during experimental acid-soss.

³ Determination of Lipide Phosphorus See p. 589.

¹¹⁰ Greenwal 1 J Biol Chem , 63, 339 (1925)

4 Determination of Total Phosphorus Dilute 1 ml of blood, plasma, or serum to 10 ml with 0 9 per cent sodium chloride solution and mix Trans fer 2 ml of this diluted solution to a 200 × 25 mm test tube, add 2 5 ml of 5 N sulfuric acid and a quartz chip, and heat over a microhurner or electric hot plate as described for the determination of acid soluble phosphorus on p 633 Foaming may be minimized by blowing clean compressed air into the digestion tube tbrough a fine tripped glass tube. When the water has been driven off and the contents of the tube turn black, continue with the digestion, oxidation, color development, and measurement as described for the determination of acid soluble phosphorus. Calculation of results is the same except that since the 2 ml of sample taken for analysis represent 0 2 ml of original sample instead of 0.4 ml as in the determination of acid soluble phosphorus, results are multiplied by 2 to give the total phosphorus, in mg per 100 ml

Determination of Phosphorus in Tissues and Other Biological Material The colorimetric method described above may be used for tissues and other biological material Dry ashing may sometimes he necessary If hittle organic matter is present, ignition with a small amount of sodium carbonate will suffice. In other cases magnesium intrate (I ml of 10 per cent) is satisfactory. For tissues, carbonate intrate fusion mix tures are perhaps best. Porcelain dishes may generally be used, but blanks should be run on them. Some silica will not interfere. If platinum is used, dissolve most of the ash with water before adding acid to remove the last of the material, since nitrous acid attacks platinum. Exaporate the total solution to dryness in a beaker or porcelain disb (covered with a watch glass as long as carbon dioxide is ovolved). Dissolve the residue in water, make to volume, and determine as usual

DETERMINATION OF SERUM PHOSPHATASE ACTIVITY

Introduction. Normal blood serum contains several enzymes or groups of enzymes which catalyze the liberation of inorganie phosphate from phosphate esters such as glycerophosphate phenylphosphate, etc. The most active phosphatase, and the one longest recognized has an optimum pH of approximately 9, and is now known as "alkaline" phosphatase to distinguish it from a second or "acid" phosphatase which is of limited activity in normal scrum but whose activity is of significance in certain pathological conditions and which has an optimum pH of approximately 5. Neither type of enzyme has any significant activity at the optimum pH of the other, although both are active at the pH of normal blood. The curzymatic activity of serum or plasma with respect to either or both enzymes is established in terms of the rate of hydrolysis of suitable phosphate ester substrates buffered to the proper pH.

Various methods have been proposed for the determination of phosphatase activity of serum For "alkaline" phosphatase, the method of Bodansky": has perhaps found the widest application. In this method, the phosphate hberated on membation of serum with buffered gly cerophosphate at pH 8 6 is used as an index of phosphatase activity, one Bodansky.

^{* 1} Bodansky J Bod Chem 99 197 (1932) 101, 93 (1933) This method is described in the eleventh edition of this book.

unit corresponding to the liberation of 1 mg. of inorganie phosphate per 100 ml. of scrum during a one-hour period of ineubation under these conditions. In the Bodansky method, the liberated phosphate is determined with stannous chloride as reducing agent, and various corrections for the influence of protein precipitant and substrate on the phosphate determination are necessary; these have been established by the author. Shinowara, Jones, and Reinhart²²² claim that these correction factors become unnecessary under the proper conditions and likewise feel that "alkaline" phosphatase activity should be determined at pH 93, the optimum pH for this enzyme. They have described procedures for the determination of inorganic phosphate. "alkaline" phosphatase, and "acid" phosphatase on as little as 006 ml. of serum. King and Armstrong 222 proposed the determination of "alkaline" nhosphatase activity in terms of the phenol liberated on incubation with buffered phenylphophate, and Gutman and Gutman224 have applied this procedure to the determination of "acid" phosphatase. A disadvantage of the King and Armstrong procedure is that the serum inorganic phosphate content, which is usually of importance along with phosphatase activity, must be determined separately with an entirely different set of reagents. In the procedures described here, the incubation procedure of the Bodansky method is used for "alkaline" phosphatase determination, with modification to permit the use of the method of Fiske and SuhhaRow (see p. 630) for the determination of phosphate liberated; for "acid" phosphatase, the conditions prescribed by Shinowara, Jones, and Reinhart are followed, likewise modified to permit the use of the Fiske and SuhbaRow phosphate method.

1. Determination of "Alkaline" Phosphatase."

Procedure. Collect about 5 ml. of whole blood in a centrifuge tube, allow to clot at room temperature, remove clot, then centrifuge (twice if necessary). The separated serum may be kept for several hours in the refrigerator, or for several days if frozen.

²²³ Shinowara, Jones, and Rembart J Biol Chem. 142, 921 (1942)

²¹³ King and Armstrong Can Med. Accor J., 31, 376 (1934)

³³³ King and Armstrong Con Med. Assoc J. 31, 375 (1934)
343 Gutnan and Gutman J. Bod Chen. 132, 201 (1940)
344 Gutnan and Gutnan J. Bod Chen. 132, 201 (1940)
345 Gutnan Gutnan J. Bod Chen. 132, 201 (1940)
346 Gutnan Gutnan J. Gutnan Gutn stoppered pyrer bottle containing an inch layer of petroleum ether Keep in the refrigerator When multiples of 100 ml. are prepared, it is advisable to distribute the substrate into small

bottles. 30 Per Cent Truchloroacetic Acid Dissolve 30 g of reagent-grade truchloroacetic acid in water and dilute to 100 ml Stable indefinitely To prepare = 10 per cent solution from this dilute I volume with 2 volumes of water To prepare a 5 per cent solution, dilute 1 volume with 5 volumes of water

Standard Phosphate Solution Place 6 25 ml of the standard phosphate solution used for the determination of imorganic phosphate (see p. 631) containing 0.5 mg of phosphorus in a 100-inl volumetric flask. Add 16.7 ml of 30 per cent trichloroacetic acid solution, and 100 ml with water acid solution. to 100 ml with water, and ms. This solution contains 0.04 mg, of phosphorus in 8 ml, in 5 per cent trichloroacetic and It should be stable indefinitely if stored in the cold Mobilette I Amnonación. Molybdate II Aminonaphtholaulfonse Acid Reagent See p 631.

- (a) INCUBATED SAMPLE Measure 9 ml of "alkaline phosphate" substrate into a glass-stoppered cylinder 11 and place m an incubator or water bath at 37° C until the fluid reaches incubator temperature Add 1 ml of serum, mlx, note the time, and incubate for exactly 1 hour Remove, cool in ice water for several minutes, and add 2 ml of 30 per cent trichloroacctic acid. Mlx let stand a few minutes, and filter through a low ash filter paper
- (b) CONTROL SAMPLE At or near the end of the incubation period, prepare a control sample as follows Measure 9 ml of substrate into a glass stoppered cylinder and add 2 ml of 30 per cent trichloroacetic acid. With mixing add 1 ml of serum, stopper, shake, and filter as above.

When both filtrates are ready, transfer 8 ml of each to cylinders or test tubes graduated at 10 ml in a third similar container place 8 ml of standard phosphate solution, containing 0.04 mg of phosphorus For photometric measurement, a fourth or blank tube is necessary. This contains 8 ml of 5 per cent trichloroaccitic acid alone. When all the tubes are ready, add to each 1 ml of Molybdate 11 reagent, and mix. Add 0.4 ml of aminonaphtholsulfonic acid reagent to each, dilute immediately to 10 ml with water, and mix. Allow to stand 5 minutes for color development.

CALGULATION For colorimetric measurement compare both control and incubated unknowns against the standard Since the 8 ml of filtrats represent two-thirds of the total sample, the calculation is in follows

Reading of Standard $\times 0.04 \times \frac{3}{2} \times 100$ = mg morganic phosphate per 100 ml serum (control or incubated)

For photometric measurement read the unknowns and standard in a photometer which is set to zero density with the blank using the same conditions as for the determination of morganic phosphate described on p 631. The calculation is as follows

Density of Unknown No. 2 × 004 × 3/2 × 100 = mg inorganic phosphate per 100 ml serum (control or incubated)

The phospi atase activity is the difference between the inorganic phosphato content of the incubrited and control samples expressed in ing of phosphorus per 100 ml. Thus if the control result is 40 mg per cent and the incubated result is 85 mg per cent the phosphatase activity is 85 - 40 = 4 a Bodansky units per 100 ml. serum

In colorumetric measurement with the standard set at 20 mm, the control tube usually reads approximately 30 mm, and a phosphatase activity up to about 8 units may be accurately determined. In photometric measurement in a 1-cm, cuvette the limit is about 12 units. For activities greater than this, the phosphate analysis is repeated on a smaller portion of filtrate from the incubated serum making up to 8 ml volume with 5 per cent trichloroacetic acid and correcting the calculations accordingly. If the hiberated phosphate is 60 mg, per cent or higher inhibition of hij drolysis occurs and the incubation is repeated with the time shortened to 30 or 15 minutes in which case results are multiplied by 18 or 33 respectively. The result on the control sample is essentially an analysis for seriam morganic pho-late but includes any effect of gly ecrophosphate on color development. This effect is frequently very low or medial, bits in which case the results on the control seriam may be accepted as the measure of the serium inorganic phosphatase values.

Interpretation. The serum "alkaline" pho plant is activity in normal adults ranges from I 5 to 40 units per 100 ml (average 2.7) and in normal

²²⁸ Or i pary test tubes and clean rul ber stor pers may be used

children 5 to 12 units (average 8 0) These values are greatly exceeded in polyostotic Paget's disease (up to 50 times), nickets (up to 20 times), hyperparathyroidism (10 times) Smaller increases are observed in a number of other diseases of osseous origin. High serum phosphatases is a main festation of processes that cause rapid growth of bone in the normal young, of new hone (repair), and of calcified and uncalcified pathological bone. Among nonosseous conditions in which increased phosphatae activity is observed are acute catarrhal jaundice and other cases of liver involvement. Under certain conditions the serum phosphatase activity may be used in the differential diagnosis of liver disease.

2. Determination of "Acid" Phosphatose " The procedure is exactly the same as for "alkaline" phosphatase, except that the buffered "acid phosphate" substrate of Shinowara, Jones, and Reinhart is used for the incubation and for the control sample Calculation of results is the same as for "alkaline" phosphatase, the unit of "acid" phosphatase activity being dined as equivalent to the fiberation of I mg per cent of inorganic phosphate during I hour incubation at pit 5 0

Interpretation. In terms of the unit above defined, normal serum entains from 0 0 to 1 1 units of acid phosphatase activity, with no significant elevation observed except in cases of carcinoma of the prostate with metastases, when values as high as 30 units or more have been observed. It should be pointed out here that the King and Armstrong unit as used by Gutman and Gutman¹²⁴ is defined in terms of the hiberation of 1 mg per cent of phenol, and is approximately double the value of the phosphate unit here described Results by the method described here will therefore tend to be about half as high as those obtained by the use of phenylpho-phate as substrato

DETERMINATION OF CHOLINESTERASE ACTIVITY

Introduction. Enzymes which eatalyze the hydrolysis of acetylcholne are known as cholinesterases and are found in both the red cells and the plasma of the blood, as well as in most other body tissues. They are particularly active in nervous tissue, as discussed on p. 297 Available vidence indicates that cholinesterase is not a single cnayme but rather anime for a group of enzymes which can be further characterized as true cholinesterase and pseudocholinesterase in terms of tissue location, physiological function, and substrate specificity. The red blood cell enzyme is considered to be a true cholinesterase, as is the brain enzyme, since its found that these enzymes act more rapidly on acetylcholine than on other choline esters such as butyrylcholine and benzoylcholine. The

^{****} Reagents Required **Acad Phorphote Substrate This is identical with the sphorphate substrate already described except that sufficient motive and is incorporated bring the pil to 6 0 into 3 100-ml volumetric flaks introduce successively 3 ml orbital learn effect and in the substrate and 5 ml of water 0.5 g, of sodium \$\textit{Region}\$-giverophosphate 0.423 g, water to describe the first flat and the substrate and 5 ml of 1 N section and Disselve by mining and water to first flat and the substrate and best flat and the substrate and the substrate and the substrate and the substrate and the substrate and the substrate and the substrate and the substrate and the substrate and the substrate an

plasma enzyme is a pseudocholmesterase in the sense that it will act rapidly not only on acetylcholme but also on a variety of other esters. Since the red cell and plasma enzymes differ in physiological function (see discussion below) as well as in substrate specificity, determination of whole-blood cholmesterase activity is relatively meaningless, the analysis being carried out on the separated red cells of plasma as required.

Estimation of cholinesterase activity is based on measurement of the rate of hydrolysis of the substrate (usually acetylcholine) under specified conditions. When acetylcholine is hydrolyzed, acetic acid is formed, and measurement of rate of enzyme action is conveniently made in terms of rate of acid production. The standard method for measuring cholinesterase activity is that of Ammon,228 based upon the use of the Warburg apparatus (see p 334), for the manometrie measurement of CO2 liber ated from bicarbonate by the enzymatic production of acid A variety of simpler methods based upon acid production have been described, of which the method of Michel, 229 described here, is believed to be one of the most satisfactory. A colorimetric procedure based upon the determination of acetylcholme present before and after treatment with the enzyme has been described by Hestrin 230 It is not always possible to compare cholinesterase activities obtained by various methods because of the sensitivity of the enzyme to temperature, pH, and particularly substrate concentration, and these conditions are rarely the same from one method to another

Method of Michel **Principle* The enzyme in an aliquot of diluted plasma or red cell hemolyzate is allowed to act on acetylcholme in a standard buffer solution for a measured at the (usually 1-2 hours) The pH of the mixture is measured at the begin ning and at the end of this time. The action of the enzyme on the substrate produces acid which lowers the pH of the mixture—the rate of change in pH is therefore a measure of enzymatic activity.

Procedure *** (a) Separation of Red Cells and Plasma. Place approximately 5 ml of freshly drawn blood, heparanized to prevent clotting, is a graduated contribuge tube and centrifuge at 2,000 r p m for 15 minutes Remove the

²¹⁸ Ammon Arch f d. ges Physiol 233 57 (1933)

Vlichel J Lab Clin Med 34 1564 (1949)
 Ilestrin J Biol Chem 180 249 (1949)

³¹¹ Reagents Required Buffer Solution I (for Red Cells) Dissolve 4 12 g soshum barbatal 054 g kllrPO; and 44 7 g KCl in about 990 ml of distilled water 4dd 28 0 ml 0 1 N hydrochloric acid solution with mixing dilute to 1 liter and mix Cleck the pill which should be 8 10 at 25°C if necessary adjust to the proper pill by tle careful furtier addition 0 tl N said or alkali as required 4dd a few drops of tokene and keep in it is refrigerator

Buffer Solution II (for Plasma) D ssolve 121 g sodium barbital 017 g KHiPO and 1754 g NaCl in about 900 ml distilled water 4dd 11 and 01 \ by drockhora acid with mixing dilute to 1 liter and mix The pli should be 8 00 at 25°C if not adjust as described above Preserve with toluene and keep in the refragerator

icetylcholine Substrate (0.11 M) for Red Cells Dissolve 2.00 g of high-grade acetyl choline chlorido in 100 ml of distilled water. Add toluene as preservative and keep in the refugerate.

Accipicholine Substrate (0 100 M) for Plasma Dissolve 3 00 g acetylcholine chloride in 100 ml of distilled water Pre-cre with folicens and store in the cold Suporin Solution 0 01 per cent supomin in distilled water

Both Buffer Solution I and Buffer Solution II show a slow decrease in pH and buffer capacity after standing for several weeks and should be checked with a pH meter before use If the pH₁ reading in a detert mation is more ti an 0.03 pH units below 8.00 a fresh

aupernatant plasma with a capillary pipet and act it aside if desired for analysis as described under (c) below Add 2 to 3 volumes of 0 9 per cent \aCl Solution to the red cells, nix well, and again centrifuge for 15 minutes Discard the supernatant saline and repeat the washing procedure, this time centrifuging for 20 minutes Note the volume of packed red cells, and remote the supernatant saline to a point where the volume of cells and saline is twice the volume of cells alon. \lik the cells well in the remaining saline, thus obtaining a 1 to diduction of the washed cell suspension in saline Transfer 0 4 ml of this cell suspension to a test tube containing 9 6 ml of 0 01 per cent saponin solution. This hemolyzed red cell solution, of which i ml represents 0.2 ml of red cells, ls used for the procedure described under (b)

- (b) Myasurement of Red Call Agriculture for an of hemolyzed red cell solution, prepared as described above, to a small beaker in containing a mill of Buffer Solution 1. Vilx and place the beaker in a water bath at 25° C for 10 minutes, for temperature equilibration. Measure the pl1 of the mix ture, using a pl1 meter and reading to the nearest 0.01 unit. Return the beaker to the thermostat Note the time, and add 0.2 ml of 0.11 M acetylcho line solution with rapid mixing. Allow the beaker to stand in the thermostat for 1 to 1½ hours. Again measure the pl1 and note the time. Calculate results as described below.
 - (c) MEASUREMENT OF PLASMA ACTIVITY Dilute 0 2 ml of plasma to 10 ml with water, and mix Transfer 1 ml of this diluted plasma to a small beaker can taining 1 ml of Buffer Solution 11 Place in a thermostatically regulated water bath at 25°C for 10 minutes, then measure the pli as described under (b) above Note the time then add 0.2 ml of 0.165 M acetylcholine solution with rapid mixing Carry out the subsequent steps as described under (h) above

Calculation The cholmesterase activity of the sample in units of $\Delta p11~{\rm per}~h^{\rm out}$ is calculated as follows

$$\Delta p \Pi / hr = \left(\frac{p H_1 - p H_2 - b}{t} \right) f$$

where pli₁ and pli₂ are the initial and final pli readings respectively t is the time in hours between the mixing with acetylcholine and the time of reading pli₁ and b and d are correction factors obtained from the following table b being a correction for non-enzymatic hydrolysis of substrate and f correcting for the effect of pli change on enzyme activity relative to buffer capacity

The results obtained by the procedure described bere represent the cholinesterage activity for 0.02 ml of red cells or plasma expressed in units of ApH per hour. If the activity per ml of red cells or plasma is desired multiply the results by 50. To express results in terms of per cent activity relative to some normal or previous value on the same individual multiply the ApH hour value found by 100 and divide by the normal or previous ApH hour value for rex one ApH hour value for rex one ApH hour value for red cell cholinesterase activity in 12 subjects to be 0.753. If an unknown sample gave a value of 0.550 then the activity in terms of per cent of normal would be 0.550/0.753 × 100 or 7.3 per cent.

buffer solution should be prepared. Pure acety leboline chloride solutions will produce a plichange of less than 0 01 pli units when added to Buffer I or Buffer II in the proportions specified in the text and in II o absence of enzyme. Any greater change indicates decomposition of the solutions and a fresh solutions abould be prepared.

²³³ Beakers which are supplied with most commercial pH meters for determining pH on small volumes (2 5 ml) of solution are very astisfactory. It is necessary to shake the beaker containing the reaction mixture for a few seconds after immers og tl e electrodes in order to obta a rapid equilibrium.

CORRECTION FACTORS FOR CHOLINESTERASE DETERMINATION

	Red Cell Cholinesterase		Plasma Cholinesterase	
pH_2	ь	f	b	f
7 9	0 03	0 94	0 09	0 98
7.8	0 02	0 95	0.07	1.00
77	0 01	0 96	0 06	1 01
7 G	0 00	0 97	0 05	1 02
75	0 00	0 98	0 04	1 02
7 4	0 00	0 99	0 03	1 01
7 3	0 00	1 00	0 02	1 01
7 2	0 00	1 00	0 02	1 00
7 1	0 00	1 00	0 02	1 00
7.0	0 00	1 00	0 01	1 00
6.8	0 00	0 99	0 01	1 00
6.6	0 00	0 97	0 01	1 01
6.4	0.00	0 97	0 01	1 02
6 2	0 00	0 97	0 01	1 04
6 0	0 00	0 99	0 01	1 09

Interpretation. In a series of 12 normal adult males, Michel found a mean value of 0.703 ApH per hour units for plasma and a value of 0.753 for red cells, by the method described here Studies hy a variety of mothods on the red cell and plasma enzyme activities of normal adults have shown that there is no sharply defined level of cholinesterase activity which may he called "normal"; though the enzymatic activity of the blood of a particular individual does not vary significantly during health, wide variations exist between individuals in a normal group. Tho method is therefore more satisfactory when used for following changes in eholinesterase activity within a particular individual than for establishing whether enzymatic activity is normal or abnormal. The factors determining the maintenance of cholmesterase activity are not too clearly understood, particularly with regard to the red cell enzyme. The plasma enzymo appears to be similar to the other plasma proteins which originate in the liver, since it decreases during liver disease and returns to normal levels on recovery. Low plasma levels have also been reported in malnutrition222 and pernicious anemia.224 The most striking lowering of both plasma and red cell enzyme is found after treatment with the anticholinesterase drugs (DFP, physostigmine, neostigmine, etc.; see p. 298) and after exposure to certain organic insecticides such as Parathion and Schradan. Determination of blood cholmesterase levels is of significant diagnostic value in these conditions.

DETERMINATION OF SULFUR

Introduction. Of the total sulfur of whole blood, a portion is present as the inorganic sulfate ion, another portion is in the form of various nonprotein organic compounds which may be present (glutathione,

M. Cance, Proc. Roy. Soc. Med., 43, 272 (1950). See however Saunders et al.: J. Nutrition, 47, 191 (1952)
 Cline, Johnson, and Johnson. Southern Med. J., 41, 374 (1948).

ergothioneine, etc.), most of which are found chiefly in the red cells, and the remainder is represented by the sulfur-containing amino acids of the proteins present In the analysis of serum or plasma, inorganic sulfate is ordinarily determined by isolation as the benzidue salt, followed by colorimetric225 or titrimetric226 estimation of the benzidine component Nephelometric227 and gasometric228 methods for inorganic sulfate have also been described. Ethereal or conjugated sulfate (i.e., the increment in norganic sulfate produced by acid hydrolysis of the protein-free sample) appears to be present, if at all, in such small amount in human plasma as to come within the limits of error in the two analyses necessary for its estimation Significant values may be found for the plasma of species other than man Total sulfur is determined by complete oxidation of organic matter, followed by estimation of inorganic sulfate present

1, Determination of Inorganic Sulfate in Setun: (Method of Letonof and Reinhold):20 Principle. Serum is deproteinized with uranium acetate. The sulfate in the filtrate is precipitated as benzidioe sulfate, then washed, dissolved, and determined colorimetrically after treatment with sodium 8-naphthogumonegulfonate

Procedure. Six ml. of 0.4 per cent uranium acetate solution are measured into a 15-mi, centrifuge tube, 2 mi, of nonhemolyzed serum are added slowly, and, after mixing by inverting four times, the mixture is centrifuged for 10 minutes. Four mi. of clear centrifugate are measured inta another centrifuge tube (selected so that the tip will retain precipitates), One ml. of glacial acetic acid and 9 ml. of benzidine solutions are added. The tube is capped, placed in ice water for at least 1/2 hour, then centrifuded for i5 min. utes at 3000 r.p.m. The supernatant fluid is decanted and discarded, and the tube permitted to drain in an inverted position for 3 minutes. Fourteen mi. of acetone are added. The precipitate is suspended in the acetone, then again centrifuged for 15 minutes at high speed. The acctone is decanted and the tube allowed to drain 5 minutes. After the mouth of the tube has been wiped, 1 mi, of a 1 per cent solution of sodium borate in 8.1 N sodium hydroxide is added and the precipitate dissolved by stirring. (The tube may be placed in warm water at 60° C, if solution is slow.) Finally, 10 mi. of water and 1 ml. of the color reagent241 are added. The solutions are mixed and allowed to stand 5 minutes, then 2 mi. of acetone are added. At the same time, 2 standards are prepared by measuring 2 and 5 ml, of benzidine hydrochloride solu-

²¹¹ For a colorimetric method different from the one described in the text, see Cuthbert on and Tompsett Buckers J. 35, 1237 (1931) Pine Buckers J. 28, 1063 (1934) 11 Power and Wakefield J. Bol Chem. 125, 665 (1938) 11 Power and Wakefield J. Bol Chem. 125, 665 (1938) 11 Dens and Reed. J. Bol Chem. 127, 101 (1926)

²¹¹ Van Slyke Hiller, and Berthelsen J Biol Chem 74, 659 (1927)

^{···} en coyac numer, and Derindsen J Biol Chem 74, 559 (1927)

"Letonoff and Renhold J Biol Chem, 114, 147 (1936) This method is said to give
lower results than other methods because of the absence of the hydrolytic effect of seid protein precipitants

One per cent benzidine in acetone, filtered and stored in a brown bottle in the refrigor ator It should be discarded when it becomes highly colored

²⁴¹ Color Reagest Dissolve 0.15 g of pure sodium β-naphthoqumone-4-sulfonate 15 over neasons. Dissolve 0 15 g of pure sodium 3-naphthoquinone-4-milionate in 100 ml of dustilled water The solution will keep alout 2 weeks in the cold Each sample of this reagent should be tested by treating 2 ml and 4 ml of the working standard solution of behandle hydrochlorule met 4. solution of behinding hydrochlorule with the color reagent, borate, water, and accione as described lelow Account. as described lelow Acceptable preparations do not deviate from the theoretical Beer's

relationship by more than 5 per cent

tion²¹² into 2 test tubes One ml, of horate solution is added to each, followed by 8 ml, and 5 ml, of water respectively. One ml of color reagent is added and the development of color carried out as described. The unknown solutions are compared with standards in the colorimeter.

CALCULATION When colorimetric comparison is made with the 2 ml benzidine standard

 $\frac{\text{Reading of Standard}}{\text{Reading of Unknown}} \times 0.02 \times \frac{100}{1} = \text{mg sulfate sulfur per } 100\,\text{ml serum}$

For the stronger standard replace 0 02 in the above equation by 0 05

Precise photometric data for this color are not available. Since the reagent and reaction resemble that for the amino-acid determination (p. 565), satisfactory photometric measurement should be possible at 490 mµ against a blank prepared by treating 10ml of water with 1 ml of borate 1 ml of color reagent and 2 ml of acctone

Interpretation. Normal human blood serum contains approximately 1 mg of inorganic sulfate (as S) per 100 ml. In nephritis with retention, the inorganic sulfate values are increased. High values have also been reported in pylone or intestinal obstruction, in leukemia, and in diabetes

2 Determination of Total Sulfur in Biological Material (Method of Stockholm and Koch) ¹⁴³ Principle The complete oxidation of biological material for the sulfur determination is often a difficult procedure. The following method in which strong hydrogen peroxide and nitric acid are used is believed to be the most effective.

Procedure. Into a 100-ml nickel crucible (50×70 mm) containing 10 ml f 25 per cent sodium hydroxide solution, introduce 0 5 to 2 0 g of the substance. The covered crucible is then heated on the steam hath until the mass is almost dry. This requires several hours, but causes considerable decomposition of the complex substances present, so that the sulfur in particular can later be easily oxidized. In case the evaporation has proceeded too rapidly, it is best to add again 10 ml of water and to repeat the slow evaporation. To the slightly moist material 5 ml of 30 per cent $\frac{1}{2}$ - $\frac{1$

The material thus partraily oxidized is next transferred to n 300 mi Kjeldahl fask, addified with nutric acld, and concentrated over a free flame until salts begin to separate This concentrated solution is then oxidized, while boling,

¹⁴⁸ Standard Benzuhne Hydrochloride Solution 0 1606g of benzidino hydrochloride puri fied as described below is transferred to n 200 ml volumetric flask dissolved in water previously warmed to about 50°C cooled and diluted to volume The solution should be stored in the cold. Ten ml are equivalent to f 0 mg of sulfur. For a working standard, 10 ml of this solution are diluted to 100 ml with water. One ml contains benzidine equivalent to 001 mg of sulfur. The solution should be stored in the cold.

The beamdon bydrochloride is purified as follows 5 g of beamdine by drochloride are dissolved in 200 ml of a per cent hydrocloric acid hydrochloric acid hydrochloric acid hydrochloric acid acid with continuous stirring. The solution is cooled in ace water for about 30 minutes when the crystals that have formed are collected on a Buchner funnel. The material is washed with cold thitted bydrochloric acid (15 ml) of concentrated acid to 100 ml of water). Uter removing the hydrochloric acid by suction the crystals are washed with work of the hydrochloric acid by suction to express a removing the hydrochloric acid by suction the crystals are washed with two 2-ml por tions of cold ethy i alcohol and four portions of ether. Meer all traces of ether are removed the dry crystals are transferred to a known bottle.

¹¹³ Stockholm and Noch J Im Chem Soc 45 19-3 (1923) 114 Merck's Superoxel

be based upon knowledge of the scrum phosphate and protein values, unless these are known to be normal

2. Method of Roe and Kahn. 252 Principle. Csienum is precentated from the protein free serum filtrate as tricalcium pho-phate, which is then determined coornnetrically by a procedure similar to that used for the determination of blood is organic phosphate (p. 630). Kuttur and Cohen have described a micromodification using this principle, in connection with their method for phosphorus determination.

Procedure.²⁴ To 8 ml of 10 per cent trichloroacetic acid in a smail flask, ^{add} 2 ml of scrum, Mix well by shaking. Filter through a Ca-free filter paper (Whatman No. 42 is satisfactory). Transfer 5 ml. of filtrate to a 15-ml conical graduated centrifuge tube, add 1 ml. of 25 per cent Ca-free sodium hydroxide solution, mix by tapping, and let stand 5 minutes. Add 1 ml. of 5 per cent trisodium phosphate solution, mix thoroughly by tapping and set aside for I hour. Centrifuge for 2 minutes. Decant supernatant liquid. place tube in an inverted position in a small beaker containing a mat of clean gauze or filter paper, and allow to drain 2 minutes. Wipe mouth of tube dry with a clean cloth. Add from a bulb pipet with a fine tip 5 ml. of alkaline. alcoholic wash reagent, forcing the wash fluid in at first so as to break up the mat of tricaicium phosphate, and then washing down the sides of the tube If necessary, use a stirring rod to break up the precipitate, rinsing down the rod with a little of the wash fluid. Centrifuge 2 minutes, decant, drain, 20d wipe the mouth of the tube as before. Repeat the washing procedute, centrifuging, etc , with a second 5-mi, portion of wash reagent, centrifuging, decanting, draining, and wiping off excess fluid as above. To the residue in the centrifuge tube add 1 ml. of acld molybdate reagent and tap against the palm of the hand to effect complete disintegration and solution of the ptecipitate. When dissolved, add 10 ml. of water, mix by tapping, and set aside Prepare a standard by transferring 10 ml of the standard phosphate solution. containing phosphate equivalent to 0 1 mg of calcium, to a second similar graduated tube, add I ml. of the acid molybdate reagent, and mix. For photometric measurement, prepare a blank tube containing 10 ml. of water and 1 ml. of acid molybdate. When all the tubes are ready, add to each 6.5

^{75, 517} Ros and Kahn J Biol Chem 81 t (1929) kuttner and Cohen J Biol Chem. 75, 517 (1927) The latter procedure as applied to the Roe and Kahn method is described in the eleventh edition of this book.

²³³ Resgents Required The trichloroacetic acid sodium hydroxide and trisodium photophate (NasPO, 12 Hol) solutions are prepared on the indicated basis from reagent the

chemicals. The two alkaline solutions should be decanted occasionally from any sodiment which forms and which might lead to errors if included in the analysis Alkaline-Alcohole Wash Reagent Str. S ml of ethyl alcohol with 10 ml. of amyl alcohol ddlines to 100 ml. of amyl alcohol with 10 ml. of amyl a

and dilute to 100 ml. with water 1dd 2 drops of 1 per cent phenolphthalem and 5 per cent Ca-free NaOll a drop at a time to a distinct pank (usually 2 to 3 drops)

Acid Molybdate Reagest Dissolve 12 s g. of reagent-grade ammonium molybdate 12 400 ml of water in a 00-ml volumetrie flask. Add slowly with shaking 100 ml of concentrated sulfurne acid. Stable indefinitely

Aminonaphtholulifonic Acid Reagent The same as described on p 631 for the determine tion of morganic phosphate

Standard Phorphate Solution Dissolve 2.265 g. of pure dry monopotassium phosphate in water dilute to I liter in a volumetric flask, and mix Add a little chloroform as a present tive This stock solution contains 0.47 mg of P per ml equivalent to 1 mg of Ca accilium phosphate The working standard is preared by diluting 1 ml of stock solution to 100 ml with water This standard about 2.4 ml of the contains 0.47 ml of the contains 1 ml of stock solutions. to 100 ml with water This standard should be prepared fresh daily 10 ml contain phosphate equivalent to 01 mg of 5. phate equivalent to 01 mg. of Ca.

²²⁴ Roe and Kahn state that only one washing is necessary. Two washings provide a greater margin of safety

ml of aminonaphtholsulfome need reagent, followed by water to the 15-ml mark, and mix immediately by inversion. Allow to stand 10 minutes before reading in the colorimeter or photometer. For photometric measurement, set the photometer to zero density at 660 mm with the blank.

CALCULATION For colorimetric measurement

Reading of Standard Reading of Unknown
$$\times 0.1 \times \frac{100}{1}$$
 = mg calcium per 100 ml scrum

The standard corresponds to 10 mg per cent scrum calcium and will be satisfactory for practically all values encountered in analyses. For very low values (below 5 mg per cent) dilute the unknown after color development to 12 ml instead of 15 ml, and correct the calculations accordingly.

For photometric measurement

$$\frac{\text{Density of Unknown}}{\text{Density of Standard}} \times 0.1 \times \frac{100}{1} = \text{mg calcium per 100 ml serium}$$

The characteristics of the color and the conditions of photometric measurement are the same as those described for the determination of inorganic phosphate (p. 632) at 660 m μ , in a 1-cm cuvette the standard has a density of approximately 0.400, and all values of serum calcium may be accurately determined under these conditions

The photometric blank will correct for any phosphate present in the color-producing reagents, but not for contamination by calcium in the earlier stages of the procedure Since it is very difficult to obtain absolutely Ca free reagents or filter paper for precise work a correction for calcium already present should be made. This is best done by running a standard calcium solution. The through the entire procedure (treatment with trichloroacetic acid filtering etc.) and comparing the value obtained with that expected. Thus if a standard equivalent to 10 mg per cent calcium gives on analysis a value of 10.5 mg per cent, the correction to be subtracted from serum results is 0.5 mg per cent. It is sometimes suggested that using such a calcium standard instead of a phosphato strudard as described, will include the calcium present in the reagents and bence serve as a correction. This is only true if the unknown and standard should happen to contain exactly the same amounts of calcium, otherwise it is in error since the constant calcium correction becomes a variable and unknown fraction of the total calcium present in the unknown.

Interpretation. See previous method

Other Methods. Numerous other methods have been proposed for the determination of serum calcium, some of which require only very small quantities of serum Instead of titrating the oxalate directly, Fiske and Adams** redissolve the washed oxalate in nitric and, dry, add oxalic and, dry again, ignite, and titrate the residue as calcium oxide Kuttner and Cohen (loc at), in a method applicable to 0 1 ml of serum, ash the material, precipitate the calcium as phosphate, and analyze for phosphate colorimetrically. Gasometric methods include measurement of the carbon dioxide evolved in the reaction between oxalate and permanganate** or ceric sulfate.**s or in the combustion of calcium pierolonate.*** the latter

³⁴⁵ Suspend 0 250 g of pure dry calcum carbonate in a little vater in a I liter volumetric flash add dilute (1 10) HCI to dissolve add a few ml of acid in excess dilute to 1 liter with water and mix. This solution is stable indefinitely and contains 0 l mg of Ca per ml. It corresponds therefore to a serum with a Ca content of 10 mg per cent.

³⁴⁵ Fiske and Valums J im Chem Sec. 53 2198 (1931)

Van Slyke and Sen Iros J Biol Chem 84 217 (1929)
 Sendroy J Biol Chem 152 557 (1944)
 Van Slyke and Kreysa J Biol Chem, 142 765 (1942)

Prior to isolation of the sodium as the triple salt indicated, the proteins are precipitated²⁴⁴ or the sample is ashed,²⁴⁷ electrodialysis may also be used for separation of the sodium from interfering material ²⁴⁵. The triple salt may be determined by graximetine,²⁴⁷ titrimetine,²⁴⁶ or colorimetine methods. Another reagent which has been proposed for the determination of sodium is pyroantimonate ^{2 o}. In the analysis of serum sodium, precision is essential because of the relatively small variations which are of clinical significance. The determination of sodium in tissues is similar to the procedures used for blood ²²¹

1. Determination by Flame Photometry. Principle The sample in solutions introduced in the form of a fine continuous spray into a nonluminous gas flame. The various ions present produce their characteristic flame spectra. By the use of proper color filters or other means the emitted light characteristic of the ion being determined is isolated and allowed to excite a photoclectric cell. The response of the photoelectric cell is measured on a suitable meter. Under controlled conditions, the meter reading is a measure of the concentration of ion being determined. A calibration curve is established by analyzing a series of standard solutions of the ion being determined. From the meter reading of the sample being analyzed, the cooccontation is established by reference to the calibration curve.

Procedure The procedure varies with different types of instrument Ordinarily a dilution of 1 230 is used for sodium analyses, i.e., i mi of plasma or serum is diluted to 250 ml with water if an internal standard, e.g., a lithium salt, is required for the successful operation of the photometer, the required aliquot of a stock solution of such a substance is added to the voluments flask before diluting to the mark. The solution is then analyzed according to the directions which accompany the instrument. In order to get the sodium content of the sample, the reading obtained is referred to a calibration currence of the sample, the reading obtained is referred to a calibration currence of the sample, the reading obtained is referred to a calibration currence.

STANDARD. Standard solutions covering the range from 100 to 150 milliequivalents of sodium per liter, at a ddution of 1.2.0 are usually adequate for calibration purposes. Prepare a stock standard containing 100 milliequivalents of sodium per liter by dissolving 5.85 g of \aCl in water and dduting to 1 liter Working standards are prepared from this stock standard by neasuring out 4.0.50 and 6 of mil aliquois into 1 liter volumetric flasks and dduting to the mark with water. These standards represent repercincy 100 125 and 1.0 milliequivalents of sodium per liter at 1.2.0 dduttion. If the analytical procedure requires an internal standard such as lithium sulfate solution, add to each working standard 4 times the volume of intimus sulfate solution rotution, add to each working standard are serum, before diluting to

²⁴⁴ Weinbach J Biol Chem 118 95 (1935) Ball and Sadusk vind., 113, 661 (1936).
Dregus Biochem Z 191 69 (1939)

DIFFERM DIFFERM 12 143 (9) (1939)

"Butler and Tuthill J But Chem 13 171 (1931) Barber and Kolthoff J Am. Chem
50c 50 1623 (1928) 51, 2233 (1929) An improved version of the Butler and Tull—
method has been described by Pisha and Spearer Arch Bioch and Bioph, a 37, 253 (1924).

"Market J Biol Chem 114, 442 (1936) Consularia and Talbott Biol. 134, 7.53 (1940).

Sobel Kraus ani Kramer ibol 116 dol (1941)

10 McCance and blupp Bookem J 25 449 (1931) Saint J Biol Chem. 96 God (1932)

10 McCance and blupp Bookem J 25 449 (1931) Saint J Biol Chem. 96 God (1932)

11 Namer and Tisdall J Biol Chem. 447 (1921) Kramer and Gittleman ibod. 6h.

233 (1924)

III For a micro-procedure applicable to small amounts of lissues and blood see Clark. Levitan Gleason and Greenberg J Biol Chem. 145, 82 (1942)

the mark and mixing. To prepare a calibration curve, read each of these standards in the photometer, plot readings against equivalent softum content in milliequivalents per liter, and draw a smooth curve to include the nomits.

Standards should always be run both before and after analyzing a series of unknown solutions, to be sure that the calibration is constant, if there is considerable fluctuation, each unknown solution should be followed by a standard closely counvalent to it

Routine precautions in the use of flame photometers include the use of gas and air for the flame which are as free from contaminating ions as possible, maintenance of a constant gas and air pressure installation of the photometer in a room free from air-borne contamination, particularly tohaeco smoke and continual checking of the zero setting and standard readings. Cleaning of the atomizer and flame chimney may also be necessary from time to time

Interpretation. See under 2, below

2 Method of Weinbach 264 Principle After deproteinization the sodium to the filtrate is precipitated in alcoholic medium as the triple salt uranyl zine sodium acctate, which is washed dissolved in water and titrated with standard sodium hydroxide. It is claimed that phosphates present do not interfere. For a similar method to which phosphates are first removed, see Dregus 265.

Procedure 373 To 1 volume of whole blood or cells for 2 volumes of serum or plasma) in a small flask, add 7 (or 6) volumes of water, shake, and let stand until hemolysis is complete. Add rapidly, with shaking, 2 volumes of 20 per cent trichloroacetic acid. making a total of 10 volumes (This deproteinization may be done with as little as 0 1 to 0 2 ml of material) Vix. let stand 10 minutes, and filter through ashless filter paper (or centrifude if the sample is small) Transfer 1 ml of the whole blood filtrate (or 0 5 ml of the serum filtrate) to a 15 ml centrifuge tube and add 5 ml of the uranyl zlnc acetate reagent From a 1-ml graduated pipet add 0 3 ml of 95 per cent alcohol and let stand for 5 minutes Again add 0 3 ml of alcohol and let stand for a few minutes. This procedure is repeated, without greatly disturbing the precipitate, until 21 ml of alcohol have been added, the entire process of precipitation taking about 16 hour Centrifuge, decant, drain by inversion on a pad of filter paper, and wipe the mouth of the tube with a cloth Wash the precipitate once hy blowing in 10 ml of acetone wash reagent, centrifuge, decant, drain on filter paper, and wipe the mouth of the tube The precipitate, which is readily soluble in water, is then transferred quantitatively to a 100-ml Erlenmeyer flash by blowing in three or four 5 ml portions of water, which has been recently bolled and cooled, to drive off dissolved carbon dlox-

¹⁷¹ Reagents Required Uranyi Zine Accide Reagent Solution 1 77 g, uranyl accidate UO(CIL(COO)) 2H O and 14 ml of glacial accts and are dissolved in about 400 ml of water by surring and heating on a steam bath and diluted to 500 ml in a volumetric flask. Solution B 231 g of zine accetate (3H₂O) and 7 ml of glacial acctic acid are blackuse dissolved and made up to 500 ml. Tha two solutions are mixed while hot allowed to stand at least 24 hours and filtered.

Actione Wash Rieggest A small amount of the triple salt uranyl zinc sodium acetate is prepared by adding 15 ml of the uranyl zina acetate reagent to 1 ml of approximately 5 per cent NaCl with subsequent addition of about 5 ml of 00 per cent alcohol in small portions bilter with suction and wash the precipitate with 4 or 5 small portions of 00 per cent alcohol and then with 4 or 5 small portions of ether sucking 072 alter each addition of alcohol or ether didd this amount of triple salt to a liter of acetone shake left stand overnight, and filter

Star dard Sodium Solution I xactly 1 g of e p sodium chloride is dissolved in water and made up to a liter in a volunctris flask. Fach ml of this solution contains 0 393 mg of sodium

.,\

Pracedure. 112 Transfer 0.5 ml. of serum to a small clean test tube containing 7 ml. of distilled water, add I ml. af 1.5 per cent sodium tungstate, mis by tapping, and follow with 1 ml. of 2.5 per cent copper sulfate solution. Stopper and shake well; then add 0.5 ml. of 2.5 per cent silver nitrate solution. Stopper and shake again; then allaw to stand 15 ta 20 minutes. Pour onto a small dry filter (Whatman Na. 5 is recammended), returning the first portion of filtrate to the funnel to ensure obtaining a clear filtrate.

Transfer 3 ml. of the filtrate ta a clean 15-ml. graduated conical centrifuse tube, 211 and in a second similar tube place 3 ml. of the standard potassium solution, containing 0.03 mg. of potassium. Add 1 ml. of 95 per cent alcohol and i ml. of distilled water ta each tube, mix by tapping, and place in a water bath at 18° to 22° C. for five minutes. Add 2 ml. of the freship prepared and filtered silver cobaltinitrite reagent, mix by tapping, and replace is the water bath. Allaw to stand 2 hours, then centrifude for 15 minutes at 284 r.p.m. Carefully remove the supernarant fluid down to the 0.2-ml. mark with a capillary pipet. Add 7 ml. af wash reagent down the sides of the tube, sianting the tube but disturbing the precipitate as little as possible. Again centrifuge for 15 minutes, decant the supernatant fluid, invert the tube. and allow to dralo an filter paper for 5 minutes. Whee excess fluid from the mouth of the tube, and repeat the washing and draining twice more.

To the washed precipitate in the centrifuge tube, add 10 ml. of 0.2 normal sodium hydroxide, breaking up the precipitate by blowing the alkali in or by tapping the tube. Place in a boiling water bath for 10 minutes. Remot, cool, and make up to 10 ml. with water. Mix well and centrifuge. Transfer 2 ml. of supernatant to a 100-ml. volumetric flask. Add 5 ml. of water, followed by i ml. of 50 per cent hydrochloric acid. and 2 ml. of 0.5 per cent sulfanilamide solution. Mix hy lateral shaking, allow to stand 3 minutes then add I ml. of the naphthylethylenediamine reagent. Dilute with water ta the 100-ml. mark, and allow to stand 5 minutes before reading in the

¹¹³ Reagents Required The sodium tungstate copper sulfate (Cu5O, 511-O), and sire nstrate solutions are prepared on the indicated basis from reagent-grade chemicals. Are? the olver nitrate solution in a brown bottle

Standard Potassium Solution Prepaie a stock standard by dissolving 2 229 g of Page dry potassium sulfate in water in a I-liter volumetric flask dilute with water to the make and mix. Preserve with a little toluene. This solution contains I mg of potassium per call and the machine machine in the machine machine in the machine in and is stable indefinitely. Prepare a working standard fresh daily by diluting 1 ml of stock standard to 100 ml with water in a volumetric flask. This solution contains 0.03 mg of potassium in 3 ml.

Silver Cobalinatrie Reagest Prepare a sodium cobaltimitrie solution as follows Dissolve 20g of crystalline cobalt nutrate in .0 ml of water and add 12 o ml of acetic and .78 (B) Dissolve 120 g, of reagent-grade sodium mirrite in 180 ml of water and add 12 o ml of scette of B to all of A, videou in the hord and if A to all of A, videou in the hord and if A to all of A, videou in the hord and if a to all of A, videou in the hord and if a to all of A. to all of A. place in the lood and blow are through the solution until all of the nitrous and fumes have been driven of This solution is stable for about 1 month if stored in the ringerator I must be filtered and the contract of the solution is stable for about 1 month if stored in the ringerator I must be filtered and the ringer and the solution is stable for about 1 month if stored in the ringer and the solution is stable for about 1 month if stored in the ringer and frigerator it must be filtered each time before using. To prepare the silver cobaltiments reagent add I ml of 40 per cent silver nitrate solution to 20 ml of filtered sodium cobaltum trate solution, shake well and filter Prepare fresh for each series of analyses.

Wash Regent Mix 2 volumes of 95 per cent ethyl alcohol with 1 volume of ether and volumes of water 2 volumes of water

⁵⁰ Per Cent Hydrochloric Acid Mix 1 volume of water and 1 volume of concentra ed drochloric acid.

upurcunore acid.

Sulfordiamité Solution Dissoère 0.5 g. of pure sulfanillamide powder (not the tablets) is a mixture of 30 ind of placest acette seed and 70 ind of water Prepare fresh weekly not hophthightendeamine Regions Dissoère 0.1 g. of N-4-in-pathible-thell lenedlamine dr. Vaphdightendeamine Bregorie Dissoère 0.1 g. of N-4-in-pathible-thell lenedlamine dr. Of the N-4-in-pathible-thell lenedlamine dr. Of the N-4-in-pathible-thell lenedlamine dr. Of the N-4-in-pathible-thell lenedlamine dr. Of the N-4-in-pathible l

mi Tubes should be cleaned with hichromate-sulfurie and cleaning mixture and rined ell with distilled water until the control of the control well with distilled water just before using. The presence of eren traces of ammons must be avoided since ammons likewise forms an used ble collationistic.

colorimeter or photometer. For photometric measurement, set the photometer to zern density at 520 m_s with a blank prepared by treating 2 ml of water in a 100-ml. valumetric flask by the pracedure described above for 2 ml. of supernatant from the alkali treatment.

CALCULATION For colorimetric measurement

$$\frac{\text{Reading of Standard}}{\text{Reading of Unknown}} \times 0.03 \times \frac{10}{3} \times \frac{100}{0.5} = \text{mg pntassium per 100 ml serum}$$

The standard corresponds to a serum potassium of 20 mg per cent, cinvering the range from 10 to 40 mg per cent satisfacturily To convert mg per cent potassium into milliequivalents per liter, divide by 39

For photometric measurement

Density of Unknown Density of Standard
$$\times 0.03 \times \frac{10}{3} \times \frac{100}{0.5} = \text{mg}$$
 potassium per 100 ml serum

At 520 m μ , and in a 1-cm cuvette, the standard has a density of approximately 0 160, and the agreement with Beer's law is such that up to 35 mg per cent may be accurately determined. For higher values, or for plantometric measurement at greater depth of solution, develop the color on 1-ml instead of 2-ml portions of the final standard and unknown supermatants. No change in calculations is involved

Interpretation. The potassium concentration of human blood serum or plasma is relatively constant, ranging from about 16 to 22 mg per cent, or 4 1 to 5 6 milliequivalents per liter The potassium content of red cells is from 15 to 20 times as great as that of plasma. This indicates the essential uselessness of whole blood potassium determinations, depending as they must largely on the red cell count, and also emphasizes the importance of preventing hemolysis in obtaining serum or plasma for potassium analyses The potassium content of plasma gradually increases on standing in contact with red blood cells, this is of importance in connection with the storage of whole blood in blood banks Pathologically, increased serum potassium has heen noted in acute bronehial astbma, in uremia, and in Addison's disease. Decreases have been noted at the time of seizure in familial periodic paralysis. The role of potassium in plasma is unknown, the plasma level may reflect to some extent the level in the body tissues and cells where potassium is ahundant, being in fact the chief base present. In the body tissues and cells potassium appears to be the physiological equivalent of the plasma sodium in osmotie pressure regulation and acid-base balance

DETERMINATION OF IRON

Introduction. The iron-containing compounds of biological material may be divided into two main groups, heme compounds and non-heme compounds. Heme iron is found in such substances as bemoglobin, cytochrome, catalase, etc., where it is firmly bound in organic combination with the porphy in nucleus (see Chapter 22) and does not react with iron reagents until after their time from such combination. Nonlieme iron on the other hand appears to be largely extractable from tissues by suitable me ins and can be determined directly in the extracts as morganic iron. There appears to be a rough correlation between the extractable iron of

No sulfonamides are found in blood or urine unless they have been administered, the level attained depends upon such factors as the size and rate of administration of the dose, the body weight of the individual, and the rates of absorption and chimination. The therapeutically effective level depends upon both the disease and the drug, blood analysis permits definition of blood sulfonamide concentration so that a given level may be altered or maintained as conditions warrant A variable nortion of the total blood sulfonamide, the exact amount depending largely upon the type of sulfonamide being used, is usually present as the acetylated derivative249 (e.g., acetylsulfanilamide) Acetylation involves the amino group attached to the benzene ring Marked differences exist between the free and the acetylated compound in respect to both therapeutic and toxic properties, distinction is therefore important. Acctylation appears to occur largely in the liver Other metabolie derivatives of certain of the sulfonamides are known, such as hydroxy derivatives, possibly combined with glucuronic acid. 200 the significance of these is somewhat obscure

Determination of Sulfonamides (Method of Bratton and Marshall) " Principle Blood is deproteinized with trichloroacetic acid. The protein free filtrate is treated with nitrous and to diazotize any free sulfonamide present excess nitrous acid is destroyed and the diazotized sulfonamide is coupled with \ (I naphthyl) ethylenediamine to form a stable red color which is then compared with that developed in a standard treated in the same way. The procedure is the same for all the common sulfonamides which have a diazotizable amino group in the molecule, the only differ ence being that a standard containing the particular sulfonamide being determined used in each case Since acctylation renders the sulfonamide incapable of diazotization only the unacetylated sulfonamide will react in the colorimetric procedure. To delet mine acetylated sulfonamide total sulfonamide is determined after acid hydrolyals which frees the amino group The difference between free and total sulfonamide represents acetylated sulfonamide

Procedure 101 Measure 2 ml of oxalated blood into a small flask and from 2

³³ Marshall Bratton and Latchfield Science 88 597 (1938)

²¹⁰ Scudi Science 91 485 (1940) also Scudi and Jelinek J Pharmacol 81 218 (1944)

²⁵⁹ Reagents Required Saponia Solution Dissolve 0.5 g of saponia in water and dilate

to I liter This solution is not absolutely necessary the saponin merely serving to haste hemolysis of the red cells. The blood may be laked with water instead of saponin solution in which sages the duty. m which case the diluted blood is allowed to stand 15 minutes before adding the trichlo acetic acid

¹⁵ Per Cent Truchloroacetic Acid Dissolve 150 g of reagent-grade truchloroacetic acid it water and dilute to I liter Stable indefinitely To prepare a 3 per cent solution from this dilute I volume w th 4 volumes of water

Sodium Aurue Solution Dissolve 6 1 g of pure sodium mitrite in water and dilute to 100 ml Prepare fresh dady

Ammonium Sulfamale Solution Dissolve 0.5 g. of ammonium sulfamate in water solution 100 mt. Stable and South Solution 100 mt. dilute to 100 mi Stable indefinitely

V (1 Naphthyt)-thylened amme Solution Dissolve 01 g of \(\) (1 naphthyt)-thylened amme Solution Dissolve 01 g of \(\) (1 naphthyt)-ethylened amme bolution dismined the discoloride (obta nable from Eastman Kodak Co Rochester \(\) Y) in water and dilute to 100 ml Store and the solution of the solutio and dilute to 100 ml Store in a dark glass bottle in the cold Prepare fresh each week 4 N Hydrochlorte And Dultue 40 m lot concentrated by drochlorte and Dultue 40 m lot concentrated by drochlorte and to 100 mil mit water Mix and titrate a 5-ml port on with standard 1 N sodium hydroxide Adjust v cancilly 4 N Stable indefinitely exactly 4 > Stable indefinitely

Standards Prepare a stock standard as follows disselve in water exactly 0.1 g of the particular sulfonam de being determ ned and tra isfer with rinsings to a 1 liter volumetr flask Dilute to if e mark with water and mix. In preparing this standard use the pure pos

buret add 30 ml of saponin solution Allow to stand for several minutes, then add 8 ml of 15 per cent trichloroacetic acid solution Mix well, allow to stand 5 mlnutes, and pour onto a dry filter

Free Sulfonamide Transfer 10 ml of the protein free filtrate to a small flask or wide test tube, add 1 ml of sodium intrite solution, and mix Allow to stand 3 minutes, then add 1 ml of ammonium sulfamate solution and again mix After 2 minutes' standing, add 1 ml of the N (1 naphthyl)ethylenediamine solution and mix The color develops almost immediately and is stable for several hours if not exposed to direct sunlight Read the color in a colorimeter or photometer as described below

Total Sulfonnmide Transfer 10 ml of the protein free filtrate to a test tube or other container graduated at 10 ml and ndd 0.5 ml of 4 N hydrochloric acid Place in a boiling water bath for 1 hour, cool, and adjust the volume to 10 ml with water Treat with sodium nitrite solution, etc., just as described above for free sulfonamide.

CALCULATION 303

For colorimetric meas irement compare the color developed on the unknown with that obtained by treating a 10-ml portion of standard solution in the same manner as the unknown Since the concentration of sulfonamide in blood may vary widely several standards should be prepared and the unknown matched against the most suitable one Satisfactory standards contain 10 05 and 02 mg per cent of the sulfonamide At a blood dilution of 1 20 these standards correspond to blood levels of 20 10 and 4 mg per cent respectively it is convenient to set the 1 mg standard at 10 mm in the colorimeter the 0 s mg standard at 15 mm and the 0.2 mg standard at 20 mm

Reading of Unknown × 20 × mg per cent concentration of standard

= mg per cent free (or total) sulfonamide in blood

For photometric measurement, transfer the colored solution to a suitable container and read in a photometer at 530 mg *** Set the photometer to zero density with a blank obtained by treating a 10-ml portion of 3 per cent trichloroscetic acid with infinite, sulfamilie etc as in the treatment of the unknown From the photometric density of the unknown and the density of a known standard preferably determined at the same time calculate as follows:

Density of Uuknown
Density of Standard × 20 × mg per cent concentration of standard

= mg per cent free (or total) sulfonamide in blood

dered drug not the tablets. This standard is stable for months in the cold. To prepare dulus standards containing 10.00 and 02 mg per cent measure 10.5 and 2 n l respectively of ite atock standard into 100-ml. volumetrie flashs containing 18 ml. of 1.5 per cent in clicroscetic acid. Dulute with water to the 100 nd. mark and mix. Store in the cold and prepare fresh every few days.

With some of the sulfonamides correction factors are necessary to correct for drug lost during the precipitation of the proteins. With sulfanilar it is sulfaguandine sulfary, dime at levels less than 5 mg per cent and free sulfathazole no correction factors are needed. For sulfapyridine levels greater than 5 mg per cent multiply the results for both free and total drug by 11 to obtain the correct values For accitated sulfathazole subtract it is uncorrected free value from the microrrected total value and multiply the difference by 13.

³²⁴ Lilters or wavelength settings from 520 to 540 run gave equally satisfactory results (see Fig. 167)

The density of a standard containing $0.2~\mathrm{mg}$ per cent sulfonamide in a 1 cm cuvette is approximately $0.300~\mathrm{at}$ 530 m $_{\mu}$ (Fig. 167). This means that at this solution thickness, bloods containing up to about 10 to 12 mg per cent sulfonamide may be read satisfactorily, since agreement with Beer's law is excellent over a wide range. Higher concentrations produce colors which are too deep for precise measurement, and in such cases the determination must be repeated at a greater dilution of the blood (or of the 1.20 filtrate diluted with 3 per cent trichloroacetic acid). If a solution thickness greater than 1 cm is used in the photometer, the range of reading is proportionately reduced. If some dilution other than 20 is used this dilution should replace the 20 in the above calculations. Bratton and Marshall point out that dilutions of 1.50 or 1.100 may be employed in this procedure if photometric measurement is used, thus permitting

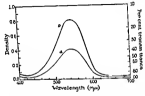


Fig. 167 Absorption Spectra of Colored Solutions Obtained in Bratton Marshall Sulfonable Method

For standards containing (a) 0.25 mg per cent and (b) 0.20 mg per cent sulfadiazine Solution depth 1 cm

the use of fingertip blood (0 1 to 0 2 mi). The only precaution is that the final concentration of trichloroacetic and in the protein free aliquot taken for analysis should be 3 per cent.

Interpretation The major factors influencing the level of blood sulfonamide content have already been presented (p. 6.8), depending upon circumstances the total blood sulfonamide concentration may range from a trace to as high as 20 mg per cent or more with a variable proportion of this in the acetylated form. It is the level of free drug which is of therapeutic importance, it is usually considered chineally that effective levels of free sulfonamide range from 3.7 mg per cent for sulfathiazide levels of free sulfonamide range from 3.7 mg per cent for sulfathiazide mas high, as 15 mg per cent for sulfamiliamide and sulfadarine. Those metabolic derivatives other than acetylated forms which have thus far been recognized do not appear to require hydrolysis before reacting in the colorimetric procedure, and will therefore be included in the value for free sulfonamide content, the extent to which this occurs, and the precise significance of these derivatives has not as yet been evaluated.

DETERMINATION OF PROTEIN-BOUND IODINE

Method of Chonev:333 Principle. The iodine-containing hormones (thyroxine and truodothyronine), along with other organic compounds containing iodine, if present, are precipitable with serum proteins by tungstic acid, zinc by drovide, etc

By digestion of the protein precipitate with chromic-sulfuric acids, the iodine is oxidized to iodic acid which remains in the acid digest. Other halides and volatile products of exidation are distilled off during the digestion

On adding an excess of reducing agents, the todic acid is reduced, and todine distilled off in a special still and collected in dilute alkali and arsenite. The concentration of rodine in the distillate is measured by its catalytic effect on the rate of reduction of coric salts by arsenious acid. This may be done by a series of colorimeter readings at 5-minute intervals, or by a continuous recording colorimeter. The method is suitable for determination of trace quantities of iodine in serum, tissue, or other biological material, in the range of 0.01 to 1 µg

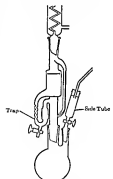
Procedure. The proteins in 3 ml, of serum or plasma are precipitated by 30 ml. of tungstic acid (1 volume 24 N sulfuric acid, 1 volume 10 per cent

sodium tungstate, 16 volumes water), or 24 ml. of zinc sulfate (1.25 per cent) and 3.0 mi. of sodium hydroxide (0.75 N) in a 50-mi. centrifuge tube, and the clear supernatant liquid is tested by the catalytic method (see below) for the presence of interfering quantities of morganic todide. If the serum contains more than 25 µg. of morganic iodine per 100 mi., additional washing of the proteln precipitate is required.

The protein precipitate is then dissolved in 20 ml, of sulfuric acid (78 per cent by weight and iodine-free) and transferred with the aid of 15 mi, additional sulfuric acid to a 300-ml, two-necked digestion flash containing 2 mi. of chromic acid (concentration 100 g./100 ml.). Digestion is carried out for a period of about 10 minutes over a 350 watt electric heater till the temperature reaches about 230° C. The digest is allowed to cool. 15 mi of water added, and the flask reheated to a temperature of 200° C.

After cooling and diluting with 20 ml. of water, the flask is attached to the special still and condenser and each of the following added through a side tube, to reduce lodic acid and the excess chromic acid in the digest, i mi. 8 \i phosphorous acid, 1 ml 0 45 N arsenious acid and 1 mi, i.5

arsenious acid, and dliuted to volume.



Iodine Still 168 IODINE DISTILLATION APPARATUS. Courtesy, Dr Albert L Chancy

per cent hydrogen peroxide. To the still trap is added 1 mi. of 1 N potassium hydroxide and distiliation is carried out for 6 minutes. The distiliate in the trap is run into a tube calibrated at 12 mi, containing 0.25 mi, of 0.15 N

³⁰ Chancy Ind Fng Chem. Anal E1 12, 179 (1940). Taurog and Chaikoff J Biol Chem 163, 313 (1940). Barker J Biol Chem. 173, 715 (1945). Connor Curtis et al. Surgery 25, 510 (1949), Kydd, Man and Peters J Clin Inrest 29, 1033 (1950), Chaney Anal Chem . 22, 939 (1950)

Chan 23

The iodine concentration in the distillate is measured by taking a 4 ml. aliquot in a colorimeter tube, adding 0.5 ml. of 0.15 N arsenious acid, and 0.5 ml. of 0.10 N ceric sulfate in 3.5 N sulfuric acid, and measuring the rate of color fading at a constant temperature (30° C.), and comparing this rate with that of known quantities of iodine (0.01-0.6 \(\rho_0\)). diluted with similar amounts of alkalı, arsenious acid, and ceric sulfate. Color measurements are made at 5-minute intervals, using a blue filter (480 m_{\rho}), and the rate of color change calculated or plotted.

Reagents should be specially selected and purified If necessary so that iodine content of quantities required for a complete determination does not exceed 0.03-0.05 µg.

Interpretation. Normal serum protein-bound iodine values range from 4 to 8 μg, per 100 ml. In hyperthyroidism, values range from 8 to 30 μg, per 100 ml, while in hypothyroid states, levels are 0 to 4 μg/100 ml Low values are also sometimes found in nephrosis and other conditions with very low serum proteins. In pregnancy a mild elevation of protein-bound iodine occurs. False high values may occur following administration of "Lipuodol," x-ray diagnostic agents ("Priodax," "Neo-Iopax," "Skiodan"), "Itrumil," and "Diodoquin."

CLINICAL MICROCHEMICAL ANALYSIS

The following discussion of clinical microchemical analysis is a brief introduction to the field, offered to the reader to encourage him to verture in and apply his ingenuity in solving problems related to the analysis of small amounts of biological fluids. Brevity precludes adequate

SCALES OF MEASUREMENT IN MICROCHEMICAL ANALYSIS, TERMINOLOGY, AND EXAMPLES IN CLINICAL CHEMISTRY

Quantity Measured Attrictions		Current Terminology	Suggested Terminology	Examples in Clinical Chemistry Currently Used					
20 Milligrams	mg.	Semimiero	Milligram chemistry	Macro-Kjeldahl analy for nitrogen					
1 Milligram*	mg.	Viero	Milligram chemistry	Micro-Kjeldahl anal)					
1 Microgram	μg, gamma, orγ	Ultramero	Vicrogram chemistry	Fluorimetric analysis for urinary coprop phryrins					
0 001 Micro- gram	пред	Submiero	Villi- microgram chemistry	Catalytic analysis protein-bound iodu					
0 000001 Microgram	μμ χ -	Vicromiero	Micro- microgram chemistry	Microbiological assay for vitamin Biz					
1 Milhliter† 1 Microliter	ml µl, lambda, or \lambda								

^{* 1} gram = 1 000 milligrams = 1 000 000 mierograms. † 1 liter = 1 000 milliliters = 1 000 009 mieroliters

recognition of the numerous original workers who did the pioneering. The definition of microchemical analysis is difficult because of the prevalence of numerous descriptive terms. The table on page 662 contains some of the terms, abbreviations, and usages The examples of analytical micro techniques used in chincal chemistry reflect the range of measurements encountered Where concentrations are high (e.g., protein in plasma = 70,000 µg/ml), minute volumes may be used for analysis, however. where low (e g, protein-bound iodine in plasma = 0 05 μg/ml), larger volumes must be used

It should be noted that many clinical chemical determinations of necessity are microtechniques, because they deal with trace amounts of material For example, quantities as small as 0.5 µg of coproporphyrin in 5 ml of urine, 0 1 μg of tryptophan in 0 02 ml serum or 0 05 μg of nodine in 1 ml serum may be concerned in a routine analysis. These figures indicate that minuteness of sample volume is not a consistent guide to the minuteness of substance measured

The advantages of microchemical procedures in clinical chemistry are numerous For example, where senal chemical data on infants are desired, standard procedures and their required volumes of blood are impractical Sobel³⁰⁶ and Natelson³⁰⁷ bave developed routine procedures requiring but a few drops of blood, readily obtained from the fingertip or beel of the infant Lowry, Bessey, and co workers developed micromethods for mass nutritional surveys, using fingertip blood for serum proteins, 308 alkaline phosphatase, 309 vitamin A, 310 vitamin C, 311 iron, 312 riboflavin,312 tocopherols,314 and thiamine 310

In addition to savings in equipment (syringes glassware reagents, and laboratory, refrigerator, and centrifuge space) it was possible to make great savings in time and perform these analyses at the commendable rate of 50 per bour Blood chemistry in small animals also requires microtechniques

Certain characteristics of microchemical analysis are uniquely favorable For example, dry ashing of 10 µl of serum takes a few minutes whereas larger amounts require considerably more time. In the analysis for calcium, if proteins are to be removed, ashing is easier than protein precipitation and collection of the supernatant fluid. In analyses using microdiffusion for separation, the rates of diffusion can be increased by reducing the reaction volumes

Some disadvantages are the special requirements of apparatus skills and methodology Kirk 116 has pointed out, however, that there is no sacrifice of accuracy in microchemistry

³²⁴ Sobel and Hanok Vikrochemie rer Vikrochim Acta 39 51 (1952)

[&]quot;Sobel and Hanok Vikrochemic rev Mikrochim 4rda 39 51 (1952)
"Natelson Am J Clin Falk 21 1153 (1943)
"Lowry and Hunter J Biol Chem 199 465 (1943)
"Lowry and Hunter J Biol Chem 194 521 (1940)
"Blessey Lowry and Brock and Lopez J Biol Chem. 164 521 (1940)
"Blessey Lowry and Brock J Biol Chem. 164 10 (1917)
"Blurch Lowry Blessey and Lowry J Biol Chem. 174 731 (1948)
"Burch Lessey and Lowry J Biol Chem. 175 475 (1918)

[&]quot;Murich Bessey and Lowin J B of Chem. 139 32 (1945)
"Burch Bessey Love and Lowin J Bud Chem. 180 1...) (194)
"Burch Bessey Love and Lowin J Bud Chem. 198 4" (195)
"Nirk Quantidative Ulrametorandigus New York John Wiley & vot 8 Inc. 1950

No new physicochemical laws apply to microcliemistry. In classical methods requiring about 1 millimole of material. 6×10^{20} molecules are measured, in microchemistry about 6 × 1015 molecules are measured, which is still a large number of molecules. This implies that reactions which take place stoichiometrically with large quantities of substance ought to work with smaller amounts, and in practice, almost all good clinical chemical procedures can be performed on smaller amounts of material Most of the facilities for measurement are limited by final concentrations of the desired substance A major device, then, in microchemistry is to limit the volume of the final reaction mixture For example, in the titration of 50 ml of 0 01 N HCl with 0 01 N NaOH using methyl red as the indicator, 100 ml of final reaction mixture is obtained at the end point With the addition of 0 05 ml (0 1 per cent of the titrating agent) a visible change in end point occurs. By reducing the volume from 100 ml to 0 1 ml, an increment of 0 1 per cent of the titrating agent should visibly affect the and point. If there is no sacrifice in accuracy of measurement, the analysis is reduced from the measurement of 500 med to 05 med with no loss in sensitivity at the small scale of analysis The same is true for colorimetry For example, 50 al of normal blood (10 ml of 1 20 filtrate) containing about 50 µg of glucose, when analyzed for glucose, will give sufficient blue color in a final volume of 25 ml to be measured in any colorimeter Reducing the reaction mixture to 1 ml and measuring the color in a suitable colorimeter allows the measurement of 2 ug of glucose (equivalent to 2 ul of blood)

Other approaches include the uso of special sensitive measurements such as fluorimetric, mierobiologic, catalytic, enzymatic, spectrophotometric, and other methods of analysis which use specific chemical, biological, or physical properties of the substance being measured The possible adaptations to microchmeal chemistry are innumerable and are facilitated by the ingenuity of the analyst and good equipment

APPARATUS AND TECHNIQUES

Precise microweighing in clinical chemistry is possible but not essential Measurements involving microprecipitates are made colorimetri eally or titrimetrically For example calcium precipitated as the phosphate is measured colorimetrically, and calcium precipitated as the oxalate 13 measured titrimetrically Wost standards are weighed on ordinary analytical balances and diluted to suitable volumes

Trace materials can usually be isolated, purified and concentrated by chromatography (resin, paper, silica gel, alumina, etc) or countercurrent distribution (see Chapter 1) Special methods may require special equip ment which can either be made by the analyst or purchased

Micropipets. Several excellent micropipets and have been described and successfully used Three are shown in Fig 169 The pipet (A) of Sisco, Cunningham, and Kirk, 218 which delivers from 1 to 1000 µl 18 widely used

III Microclemical Specialities Co. 1834 University Ave. Berkeley Calif , lists about types of micropinets assessed to be a special transfer of the control o 12 types of micropipets ranging in volume from 0 25 to 1000 µl 114 Sisco Cunningham and Kirk J Biol Chem 139 1 (1941)

and readily available commercially. The straight tube pipet (B) is simple, is easy to construct and calibrate, and can deliver accurately volumes of 1 to 1000 µl or more. C represents a self-filling pipet used for automatically measuring 1 to 10 µl. This type may also be used as an overflow pipet. The constriction type pipet made by Levy³¹⁹ is a very convenient micropipet delivering from 1 to 300 µl when properly made and used Levy;³¹⁹ Lowy,³¹⁹ and Natelson³⁰⁷ privide details of construction. As an alteruative, straight tube blowout pipets (Fig. 169, B) with fine tips may be used. When rinsed with water in reaction mixture, they will deliver contained volumes with errors in 0.1 to 0.5 per cent. Pipets of over 20 µl

when emptied by dramage alme will deliver volumes with 1 per cent error or less The error is smaller in the larger pipets Siliconed pipets deliver the contained volume with out dramage error, 120 heuce they can achieve excellent replicability down to 0 1 per cent error Turther more, the complete dramage climi nates the ueed for rinsing hetweeu samples This advantage fails with whole blood but holds with blood filtrates Pipets are calibrated to contain or to deliver, depending upon their use The delivered or contained (as desired) weight of water is a convenient hasis for ealibrating pipets The weight of water in mg × 1004 (at 25° C) gives the volume in al Mercury weight (in mg) delivered $\times 0.0739$ is the volume in all Other conven ient methods involve measuring the delivered quantity of chloride, acid, or dve from a solution of known concentration Unsiliconed pipets

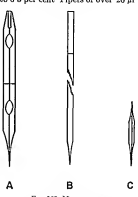


Fig 169 Micropiples.

A Pipet of Sisco Cunningham and

A Pipet of Sisco Cunningham and hark B straight tube pipet C self filling or overflow pipet depending upon how it is used

used to deliver should be emptied slowly and for the length of time specified in the ealibration, and then the remaining contents should be emptied by gently blowing while the pipet is rotated slowly to transfer the tip contents onto the wall of the container. A fine tip allows excellent replication

Microhurets There are a number of suitable, ingenious microbinets described in the literature, many of which are made commercially Especially sitisfactory is the Scholander¹ type, an example of which is shown in Fig. 170 This buret was made from a machinist's 1-inch microm-

²¹⁹ Lovy Compt rend trav lil Carlibery 8 r chim 21 101 (1936)
210 D gg u ni 15 itl Sci sci ec 116 3 bs (195...)
210 cd lander Science 95 1 - 7 (11)

etcr, which can be obtained with a scale calibrated in very small fractions of a millimeter or inch. As the micrometer is turned clockwise it propels the spindle into the mercury as shown in Fig 170 and displaces a volume equivalent to the volume of the shaft that enters the mercury, simil taneously displacing an equal volume of the standard solution above the mercury \ fine opening in the buret tip prevents back diffusion of the reaction mixture into the buret A titration is made to the desired end

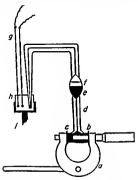


FIG 170 SCHOLANDER MICROBURET AS USED WITH THE MICROTITEATION TANDS. SHOWN IN Fig 171

The setup allown is for the electrometric determination of elloride a 1 inch microm cter and rod used for holder b gasket c set screw an I metal plate d glass luret e mer cury f carbon tetracl lori le q special clec trodes $h \to \times 1$ mm vial which holds 1 ml of reaction mixture a vial holder attached to staft of titrator

point whereupon a reading of the micrometer is made and a new titration is begun. The tip is not washed between titrations. After the burct 15 emptied more standard reagent is added by turning the micrometer handle counterclockwise The tip should be wide enough to allow rap d filling without pulling in air through the joint. The usual micrometer will deliver about 30 µl per mm (or 75 µl per 0 1 mch) and the buret will deliver this volume to ±003 µl or 01 per cent I or ordinary work one filling of standard solution will provide for 8 to 25 continuous titrations Smaller deliveres with equal precision can be achieved by reducing the diameter of the shaft which cuters the mercury

The mercury di placement huret has been criticized on three points

(1) expansion with heat, (2) leaks through couplings, and (3) reaction of mercury with reagents above it. Expansion errors are negligible when wide temperature fluctuations are avoided. Leaks are rare in properly maintained burets. The problem of reagent interaction with mercury is com-

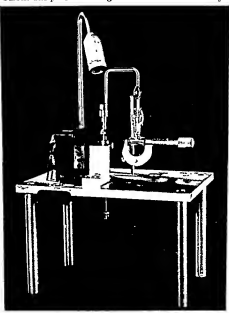


Fig. 171. Microtitration Table with Mounted Scholander Buret.

The vial fits into a holder attached to the rotator. The litter can be raised and lowered. The buret tip functions as a surring rod Courtes, Walter Reed Army Medical Center, Washington, D. C.

pletely eliminated for most aqueous reagents by interposing a 3 to 5 mm. (about 0.3 ml.) layer of carbon tetrachloride between reagent and mercury.

A microtitration table, 322 devised so that the buret described here can readily be mounted, is shown in Fig. 171. When the vessel into which the

m Made by H. Kugler, College Park, Md.

buret tip is immersed is rotated rapidly the buret tip stirs the reaction inixture Peaction vessels from 0 1 to 1 ml may be used and rotated at



Fig. 1.2 Dragray of CONTAN DIFFUSION Calz.

200 to 900 rpm Any sensitive indicator or pecial

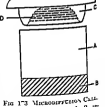
clectrodes using standard comment and circuits can be used for titrations At the end point the rotating shaft is lowered, a new titration vessel is inserted into the holder, and the shaft is raised into position for the new titration Glass vials of a uit able diameter are cut to vertical size for titration vessels A white peniculin bottle stopper223 is suita ble for holding volumes of 100 µl Acidimetric titra tions in 1 ml volumes may be performed us ng 2 standard pH meter fitted with glass quinhydrone or antimony electrodes, the latter two kinds of electrodes have been used routinely in 100 µl vol umes Istration curves of excellent quality and

accuracy are obtained The calomel half cell (see p 47) is made from a sidearm test tube with the sidearm bent and drawn out to a fine up Year is used to plug the tip which is in

serted into the difration vessel along with the opps its electrode and the buret tin

Reaction Vessels Titration vessels can be made ir m class vials cut to a suitable size Standard small test tubes of 3-ml (10 × 70 mm) or f ml (6 × .0 mm.) volume serve as reaction vessels which can be heated, cooled or otherwise manipu lated Such test tubes can also be made uto volumetric flasks Small centrifuce tubes from 0 f ml up and high speed mi crocentrifuges are available commercially

Microdiffusion Apparatus The Con way cell3 4 (1 ig 172) for diffusion has been well and universally studied and applied In conventional use, the reaction mixture is placed in the outer chamber and the receive selution in the inner chamber The dish is scaled with a lubricated class plate. Any liberated gas diffuses from the outer to the inner chamber fu the deter mination of Mi, the inner chamber con tains ifCl or other suitable seid for the determination of CO2 the inner chamber



i lox lomm val Breaction mixture which liberates gas or absorbs it C rul ber pen e llin bottle stopper D well in the stopper wich contains solit on that receives gas of I berates it In the determination of COz gas is liberated with lactic acid at B and received in Ba (OII) at D In the determ nation of ethanol blood is placed at D and the I berated ethanol oxil zedly ac d d chromate at B

cor tains Ba(OH). The determination is made by titrating the solution in the mi cr chamber or w thdrawing the contents for colorimetric analy (A) to contain the reaction mixture, and a penicillin-bottle stopper (C) for the receiver. In the determination of bicarbonate, ²²⁵ and is placed at B and a measured quantity of $Ba(OH)_2$ is placed at D. The sample (blood) is introduced into the vial and the liberated CO_2 is received by diffusion into the $Ba(OH)_2$ above. The stopper (D) is inverted and placed on the titrator and the excess $Ba(OH)_2$ is measured and imperiorally. Another example of the use of this cell is in the measurement of blood alcohol where 50 to $100 \, \mu$ l of blood are placed in stopper D and 0 500 ml of standard

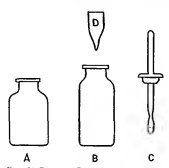


FIG. 174 DIFFUSION BOTTLES FOR THE DETER

A Pencilln type bottle B streptomy en type bottle C glass receiving rod 6 × 90 mm with a constriction 10-15 mm from the end The stippled area represents a roughened surface (ground) to aid in retaining the film of acid which receives the diffused NH, D cap for bottle B

acid dichromate in the vial at B. The diffused alcohol is measured by titrating the excess dichromate remaining after ovidation of the ethanol. Another group of diffusion cells²¹⁸ is shown in Fig. 174—1 is a standard

pencillin bottle *** C is a glass rod fitted into a pencellin bottle stopper. The rod has a constriction at one pole. The tip surface is ground so that when inserted into acid it will retain a film of the acid. A sample containing NH₂ is introduced into 4. The final volume is adjusted to 1.0 ml. The solution is made alkaline with K-CO₂ crystals, the receiving rod, previously dipped into 1. Y H-SO₂ to the constriction, is inserted into A and closed. The entire cell may be placed in a rotator (Fig. 175) and rotated so that diffusion is accelerated. The NH₂ collected on the receiv

¹²³ Seligson and Seligson Inal Chem 23, 15:7 (1951)
224 Seligson and Seligson J Lab Clin Wed 38 1.4 (1951)

²²⁷ Type 5- 05 NO-50L-VIT bottle T C Witaton Co Millydle V J

ing rod is measured colorimetrically B is a streptomycin²²⁸ bottle in which a Kieldahl determination may be carried out

Photometry. Any one of the several available photoelectric color imiters may be used for microchemistry. The Lowry and Bessey adapta tion for the Beckman spectrophotometer has the advantage that volumes down to 25 µl can be examined over a wavelength range from 235 to 935 mμ Microcuvettes (3 × 10 × 25 mm inside measurements) hold 08 ml and may be used in some of the newer instruments without

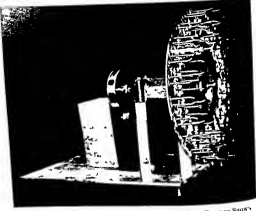


FIG 175 ROTATOR FOR FACILITATING DIFFUSION OF NH; IN BOTTLES SHOWN rs Fre 174

Courtesy Walter Reed Army Med cal Center Washington, D C

special carners. For smaller volumes a sheet of metal with a 10– $14\,\mathrm{mm}$ hole narrows the beam of light so that it will pass through 100 µl or less Lowry states that ruising between samples is often unnecessary in sensi determinations of similar samples Siliconed cuvettes eliminate the need for running

CLINICAL MICROCHEMICAL PROCEDURES

Determination of Urinary Ammonia Nitrogen 134 Principle Ammonium 1003 are converted to \11, by alkalinization the literated \11, gas is collected by diffusion and measured colored an I measured colometrically. The working range is 5 to 10 µg of NHr. It may be reduced to 0.5-1.5 µr. be south 1. red acd to 05-15 µg by suital k adjustments. This basic method is applicable to

[&]quot; Type S-281 NO-SOL-VIT bottle T C Wieston Co Millville N J " I procell Ma ufacturing Co 207 F 84th St N Y C

every system which liberates NH₁ stoichiometrically For example urease will hydrolyze urea and will hydrolyze amides (glutamine urea etc.) and ninhydrin will oxidize amino acids to liberate NH₁. The classical Lyeldahl method can also be applied as illustrated below

Procedure ¹⁵⁰ Add 50 μ l of urine to a diffusion bottle (A, Fig. 174), and water to 10 ml and 1 g k. CO₁, 13 μ H₂O. Stopper with the receiving rod (G_i , Fig. 174) previously dipped to the indentation in 1 N II SO₄. Place the bottle on the rotator (Fig. 175) and rotate for 30 minutes (complete diffusion). Transfer the rods to clean bottles or cuvettes and wash the and into the bottle or cuvette with 10 0 ml of diluted Nessler reagent. Read in 5 minutes at 420 m μ and compare with a standard (1 0 ml) and a blank (1 0 ml) of water) treated as described for the diluted urine sample.

If speed is essential, rotation may be stopped at 10 minutes and the meomplete diffusion (90 per cent) corrected by comparing unknowns to similarly run standards. If smaller amounts of Nessler reagent are used, the working range diminishes accordingly. One μg of introgen per ml of Nessler reagent is a good ratio. After diffusion the collected NH₃ on the receiving rod can be stored in a clean bottle and measured at the analyst's convenience. The error is 1–2 per cent. Where air contamination by NH₄ is a problem, equipment should be runsed and dired before use

Determination of Urea in Blood, Urine, Serum or Other Biological Fluids ***I Principle* The method above is applied by measuring the NH; liberated by the hydrolysis of urea with urease The same apparatus is used

Procedure Transfer 50 µl of serum or 10 µl of urine to a diffusion bottle, make to 10 ml with 005 Nl phosphate buffer, pH 70, and add 1 drop of fresh 5 per cent urease suspension After 15 minutes at 40°C, add 1 g K2CG, 1½H20 and cap with a stopper and recelving rod previously dipped In 1 N II SO. Rotate for 30 minutes Measure the collected NH, as described above, comparing with NH; standards and a reagent blank. Correct urine for contained NH, determined as described above. Fresh blood or serum requires no correction Measurements can be made on 5 µl of sample if desired

Analyses of a sample of urea on 8 different days produced a standard error of 0.5 per cent. All the usual precautions for protecting urease apply

Determination of Total Nitrogen by Myeldoll Digestion and Microdiffication. The sample is digested with a concentrated acid digest on mixture until all of the nitrogen present has been converted into aminonia. The ammonia is released by alkali as usual and collected by microdiffusion. The collected ammonia is determined colorimetrically. The method as desembed is for the determination of non-protein nitrogen (NPN) in a protein free plusina filtrate. It is equally npj lical le to the determination of total N in urner plasma prote is set. The size of sample is ould be such as to contain not more than approximately 16 ug of total V.

³¹⁶ Reagents Required Standard (VH.42SO4, 15 µg V per ml

^{1 \} H_1SO ... K_1CO : $I_{22}H_1O$ Crystals Fit bottle with spoon to deliver about 1 g.

Vessler Reagent of Vanselow (Ind Eng Chem Anal Ed 12, 516 (1910)) Dissolve 34 9 g KI and 45 5 g High in as little water as possible Add 112 g KOII and water to 1 liter let stat 13 days dilute 15 before using

ar 13 days unite 1 3 before using

13 bel geon and Bollier. To be published

Procedure 223 A protein free filtrate is obtained by adding 9 parts of 34 per cent trichloroacetic acid to 1 part of plasma, and filtering Add 0.5 ml of 1 10 filtrate (equivalent to 50 al of serum) in other sample to the hard glass bottle (B, Fig. 174) followed by 1 0 mi nf digestion mixture. Heat slightly on a hot plate to just short of boiling, and continue heating until all traces of water disappear from the bottle, as manifested by an absence of droplets at the sides Slowlyraise the hot plate temperature to about 500° C and add cap (D, Fig. 174) to prevent loss of spray A suitable temperature is achieved when the sulfuric acid refluxes one half way up the bottle Heat for 2 bours Cool and wash down caps and sides of flasks with 1.5 ml of alkaline washing sola tlon using a fine tipped pipet. Add 4 tn 5 pellets of NaOH with tweezers and stopper with an acid coated receiving rod (C, Fig. 174) as described on p 61 for the determination of urinary ammonia Rotate 90 minutes for complete diffusion Wash the ammonia from the rod with 10 0 ml of Nessler reagent and measure the color as described previously Prepare standards and blank 20 1151121

Determination of Chloride *** Principle This is a modification of the Cunnisham** procedure The sample of plasma urine or off of fluid is made strongly and the cl1 ride is titrated with \(\frac{1}{2} \) \text{Vol. uotil the special electrodes give off current indicating the appearance of Ag* and the disappearance of Cl

Procedure ³³³ Clean two electrodes ³³⁴ by repeated immersion in 10 per cell INOs. Amalgamate one electrode hy repeated immersion in clean mercur heneath 10 per cent IINOs. The silver electrode tip is stored in water, the amalgamated one in mercury Place 50 sl of serum or urine in a vial, said drops of 3 6 N II;50s. Place the vial on the titration assembly (Fig. 171) addition to the control of the vial on the titration assembly (Fig. 171) addition to the control of the vial and the vial should rotate at about 600-800 r pm. The whining diagram is shown in Fig. 176. Adjust the galvanometer to zero with the aid of back current from the potentiometer. Add AgNO, from the buret until the galvanometer indicator moves to and remains for 15 seconds at a predetermined point on the scale. The buret and the contained AgNO, can be calibrated against standardiorides obultion to give readings of chloride containstrated against standardiorides obultion to give readings of chloride containstrated against standardiorides obultion to give readings of chloride containstrated against standardiorides obultion to give readings of chloride content directly. The end point is chosen as the inflection point of a titration curve made by plotting givenometer unite against buret units.

The method is adaptable to one µg of chloride by reducing volumes, using smaller electrodes and more dilute solutions

Determination of Blood Alcohol Contential (Modified from Winnick 228) Principle The alcohol is separated from the blood by microdiffusion and collected in

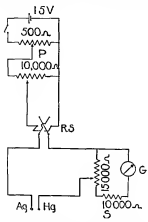


FIG 176 CIRCUIT FOR MEASUREMENT OF CHLORIOE

Ag silver wire electrode Hg silver wire elec trode amalgamated on its surface, G galvanometer sensitive to 734 × 10 10 amp (G L 713) S shunt for the galvanometer, RS reversing switch P, potentiometer made from two variable resistors and a 1.5-volt battery See Fig 171 for electrode and microburet arrangement on the microtitration table

standard chromic acid solution where it is oxidized. The chromic acid remaining is titrated indimetrically with thiosulfate. The thiosulfate is standardized by analyzing a standard alcohol solution

Procedure 239 Illood is prevented from clotting by the use of sodium fluoride, and if not used may be kept in the refrigerator for 24 hours Pipet 250 pl of

¹²¹ Seligson and Sleeman Unpul list ed

Winnick Int Fng Chem Anal Ed 14 523 (1942)

Reagents Required Standarl Chromic Acid Solution K1Cr1Or 0 05 N in 10 N H-SO4. 02 V Va-5 O

l'otage em sods le _0 per cent March Sol Ho : 0 1 per cent

kiCOi 110 IIIO C I Var dard theolol 160 rag /100 ml Dilute 2 00 ml of ab-lute alcohol (sp. g. 0.800) to 1 liter with distilled water and mix

the standard chromic acid solution into a glass vial (A, Fig. 173) Pipet 50 μ L of blood into the well of the rubber stapper (D) Add about 0 1 g K,CO: (large crystals) to the well. When this is mustened by the blood, no carbonate will fall out. Stopper the vial and allow to stand overnight in an aven at 37° C. to allow diffusion and uxidation to become complete Run a parallel deter mination on water alone (50 µl) as a blank, and on 50 µl of standard alcohol

When diffusion is complete, remove the stoppers and titrate the chromic acid in each vial as follows Add 1 drop of KI solution, place the vial on the titrating table, and titrate with standard (0 2 N) thiosulfate until the solu tion is faintly yellow Add I drup of starch solution and titrate to the disappearance of the blue color

CALCULATION Subtract the buret units required for the standard from the value found for the blank Call this value n Subtract the buret units required for the sample from the value for the blank Call this value b Since the standard represents 160 mg. per cent alcohol, the calculation is as follows

mg per cent alcohol in blood =
$$\frac{160}{a} \times b$$

The alcohol in as little as 10 µl of blood can be measured by reducing the concentration of chromic acid and thiosulfate Higher temperature will accelerate diffusion and oxidation. The error of the analysis is ap proximately 3 per cent

Microdetermination of Acid or Alkali Principle The solution to be titrate is placed in a titration assembly fitted with a suitable pair of electrodes (calomel an either glass quinhydrone, or antimony 100 electrodes) attached to a pH meter Thep. is measured after each increment of added standard alkali or seid as the case may b Titration is continued to the desired off end noint

Procedure Place 10 ml of the solution to be titrated in a 15 \times 17 mm $^{\rm stal}$ Place in the holder of the titration table (Fig. 171) and insert the electrodes, which are connected to a pli meter Start the rotating shaft and read the pli as increments of standard alkall (or acid) are added Read the buret when the desired pil is reached

With this system, titrations can be carried out rapidly to the desired end point For acid solutions approximately 0.01 \in strength, 0.2 alkali is satisfactory By reducing volumes and concentrations and em ploying antimony or quinhydrone electrodes as little as 100 µl of solution may be titrated Errors due to contamination by the CO2 of the air may be eliminated in most instances by blowing a gentle stream of COr free air over the surface of the titrating vessel

The procedure described here has been found particularly useful in microbiological assays hased on measurement of acid production Titrations were carried out in a final volume of one ml , in 15×15 mm vials which were incubated in a Petri dish modified to hold 25 vials spatula tip of quinhy drone was added to each vial contents, and titration continued to pil 70 using a platinum calomel electrode system. With one filling of alkali in the buret from 25 to 50 yials could be titrated at a rate of better than I per minute, with an accuracy exceeding the requirements of the method.

Measurement of pH and Hematocrit. Scholander342 and others bave devised accurate microgasimetric methods for CO2, N2, O2, and CO. For clinical purposes the method of Shock and Hastings (see p. 703) for pH. CO, and hematocrit is very convenient and desirable. Singer and Hastings 343 provide a helpful nomogram and discussion for evaluation of various forms of acidosis and alkalosis, using the data obtained by this method.

Measurement of Sodium and Potassium. Suitable but relatively tedious microchemical techniques have been established for the determination of sodium and potassium With the advent of flame photometry. rapid, accurate micromeasurements are easily achieved (see pp. 650 and 653). Sodium (0.7 μeq.) in 2.5 ml. of 1:500 serum (5 μl.) can be measured in most available instruments. Potassium (0.1 aeg) in 2 5 ml of 1:100 serum

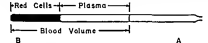


FIG. 177, COLLECTION OF BLOOD IN A MELTING-POINT TUBE.

A, Tube, 1.5-20 mm, O.D × 100 mm. The end at B is sealed and the tube is centrifuged to pack the red cells. Tho hematocrit equals mm packed red cells/mm blood, Cutting the tube at the red cell-plasma interface allows exact separation of plasma for microanalyses

(25 μ l.) can similarly be measured, both with an error of 1 to 2 per cent. Increased sensitivity can be achieved by including acetone in the diluent.

Other Methods. Natelson307 has described a system for measuring hematoerit, plasma Na, K, Cl, protein, urea and glucose ou 100 µl. of fingertip blood. Blood is drawn by capillarity into a 1.5 × 11 mm. tubes44 (ca. 0.1 ml., Fig. 177), sealed at one end, and centrifuged at 4,000 r.p.m. for 10 minutes. The fractions are measured with a rule and the volume of packed red cells calculated The tube is cut at the red cell-plasma junction and analyzed for the constituents named above. Sobel345 has described a microtitration method for ealeum which appears to be more reliable than micro flame-photometry methods. Total serum proteins and albumin-globulin measurements may be made by the bluret346 colorimetric technique except that reaction volumes are reduced to meet the need of sample size. For example, 100 µl. of serum in a final volume of 10 ml. in the standard bigret procedure can be reduced to 10 al. in a final volume of 1.0 ml. Similar reductions in any desired salting-out procedure can be made; bowever, the usual preeautions concerning denaturation of

Scholander, I'lemister, and Irving. J. Biol. Chem., 169, 75 (1947).
 Singer and Hastings. Medicine, 27, 223 (1948).

¹⁴⁴ Kimble 34500, wet with heparm and dried before use.
¹⁴⁵ Sobel and Sobel: J. Biol. Chem., 129, 721 (1939).

³⁴ Gornall, Bardawill, and David: J. Biol. Chem., 177, 751 (1919).

albumin by excessive shaking in the salt mixture and ether, and the variation due to differences in salt mixtures, should he observed Paper electrophoresis (see p 464) makes it possible to fractionate proteins in serum obtained from a few drops of fingertip blood

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(Methods for the analysis of blood for bicarbonate content, blood gases, and hydrogen ion concentration are guen in Chapter 24. Methods of analysis for various vitamins will be found in Chapter \$5, and for penicillin in Chapter \$6)

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24

Respiratory Exchange and Neutrality Regulation

General. The major function of the lungs in respiration is to facilitate the addition of oxygen to the blood and to remove excess carbon dioude from the blood. The oxygen is carried to all the tissues, where it is utilized in the metabolic processes taking place within the cells. The chief end products of these metabolic processes include H10, CO2 urea, and various organie and morganie acids such as lactic, unc. phosphone, sulfuric, and the like The production of end products which are bases, such as am monia and the various organic amines is relatively unusual or confined to lower species or to special tissues Of the various end products listed only H2O and urea are neutral substances and can be excreted by the bods without bringing into play the various mechanisms of neutrality regula tion which are the subject of this discussion

The CO2 produced in the tissues is removed by diffusion into the blood where the major portion (approximately 70 per cent) at once undergoes hydration to form carbonic acid the remainder combining with the I lood proteins (including hemoglobin) to form carbamates (20 per cent) of remaining in physical solution (10 per cent) The newly formed carlome acid must be immediately neutralized or the blood would become far more acid than is compatil le with ble, this neutralization is mediated largely through hemoglobin as described below. The other acid end products of metabolism are neutralized as soon as they are formed and exist in the I lood and tosues as salts neutralization being effected through reaction with such ions as the HPO. ion largely in the cells and the IICO, ic? largely in the blood plasma. Thus carbonic acid is not only an important end product of tissue oxidation but it also plays a significant role in neutrility regulation. The level of the blood brearbonate content is the most satisfactory single index of the ability of the body as a whole to neutralize acid end products of metabolism hence the term The peculiar virtue of the carbonic acid bicarbonate sy tem in centrelling the neutrality of the body lies in the volatility of carbon dioxide and hence its ready chammation by the lungs. The nonvolute act is (une plot phore (te) after conversion into their salts can be dis-

posed of by the kidneys Role of Oxygen V complete understanding of the mechanism of neurality regulation and CO2 transport in the Hood requires knowledge of the part played by hemoglobin in the transport of oxygen (see Fig. 178) Oxygen is found in the blood in two forms, (a) in physical solution, and (b) in combination with the respiratory pigment hemoglobin to give the compound oxyhemoglobin. The amount of oxygen in physical solution in

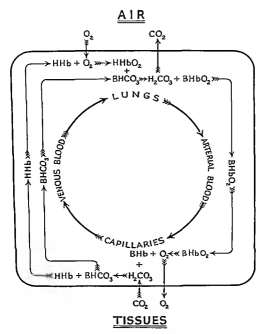


FIG. 178. THE ROLF OF OXYGEN IN THE TRANSFORT OF CO; BY THE BLOOD

the blood is relatively small, most of the oxygen present being in the combined form of oxylicmoglobin. An equilibrium exists between the free oxygen in solution, the hemoglobin which is not combined with oxygen (so-called reduced hemoglobin), and oxylicmoglobin, this relationship can be written as follows.

enters than can be neutralized in this manner the blood becomes more acid it its hydrogen ion concentration increases

To understand the nature of this change, it is necessary to consider the general properties of huffer solutions (see also Chapter 1) A buffer solution ordinarily contains a weak acid and its salt, or a weak base and its salt Such solutions are capable of taking up limited amounts of acid of base with much less change in hydrogen ion concentration than would result from the same addition of acid or base to water or to sodium chloride solution, neither of which has buffer power A buffer system can act therefore as a reservoir of alkali for the neutralization of the acid end products of metabolism The officiency of a buffer system in resisting change in reaction is greatest at the half-neutralization point, ie, when the molar concentrations of salt and acid are equal Fig 179 represents a titration curve for a weak acid (carbonie), that is, a curve in which the per cent of total acid neutralized is plotted ogainst the pH at each step in the titration The S shaped curve is characteristic of buffer acids of alkalies, and demonstrates that when of per cent of the acid is neutralized (ratio of salt to acid = 1), the slope (rate of change of pH) is least

The relation between the reaction (pH) and the ratio of buffer acid (H4) to buffer salt (BA) is derived as follows. The equotion for the electrolyte dissociation of a weak acid into hydrogen ions and anions is $HA = H^{\dagger} + A^{-}$ from the law of mass action we know that the velocity of the reversible reaction in either direction is proportional to the concentrations.

of the reacting constituents That is.

Velocity (left to right) =
$$l_1HA$$

Velocity (right to left) = $l_2(H^+ \times A^-)$

in which the symbols also represent concentrations. At equilibrium the rate of reaction in each direction is the same, the equilibrium being disame. Therefore

$$I_1IIA = I_2(II^+ \times A)$$

or transposing,

$$\frac{P_1}{I_2} = \frac{(H^+ \times \Lambda)}{H\Lambda} = K$$

K being the dissociation constant of the acid Transposing again,

$$\Pi^+=K\times\frac{\Pi\Lambda}{\Lambda}$$

In a solution of a weak acid and its salt only a very small fraction of the amon Λ , originates from the dissociation of the free acid the rest coming from the dissociation of the salt BA into B* and Λ . Most salts in the concentration found in body fluids are ionized to the extent of 60 to 90 per cent. If the degree of dissociation is represented by λ_1 the concentration of amons $\Lambda=\lambda B \Lambda$ and

$$H^+ = K \times \frac{H \Lambda}{\lambda B \Lambda}$$

Since λ varies to a relatively slight extent over rauges of concentration within such limits as are found in blood constituents, it may be stated as an approximation that

$$\frac{K}{\lambda} = K_1$$

and

$$H^+ = K_1 \frac{HA}{BA}$$

 K_1 is called the "apparent dissociation constant"

By definition pH = $-\log H^+$ (see p 30) (For example, the hydrogenion concentration [H+] at neutrality is 0 000,000,1 normal = 10^{-7} Therefore the pH = $-\log 10^{-7} = -(-7) = 7$) Taking the logarithms of both sides of the last equation,

$$\log H^+ = \log K_1 + \log \frac{HA}{BA}$$

and, changing signs,
$$-\log H^+ = -\log K_1 - \log \frac{HA}{BA}$$

Substituting pH for $-\log H^+$, and, for an analogous reason, p K_1 for $-\log K_1$,

$$pH = pK_1 + \log \frac{BA}{HA}$$

This equation is known as the Henderson-Hasselbalch equation and is a most fundamental concept in the understanding of acid base equilibrium in the body. The value of the constant pK_1 for various acids is numerically equal to the pH when the ratio of salt BA to acid HA is unity, since the log 1=0 Under this condition, it will be recalled, the maximum efficiency of the buffer action obtains. The following table gives the pK_1 values for the more important buffers in blood

Buffer System	pK_1					
ВНьО, ННьО,	7 10					
внь ннь	7 3					
BHCO, H,CO,	6 1					
B-HPO, BH-PO,	6.8					

With the Henderson-Hasselbalch equation in mind, consider what happens when acid is added to a buffer system. The added acid reacts with the buffer salt present to produce an equivalent increase in the amount of buffer acid, at the expense of the buffer salt, together with the salt of the entering acid which plays little, if any part in pH change and may be disregurded. The decrease in buffer salt concentration and corresponding increase in buffer acid concentration, however, mist necessarily change the pH of the solution in accordance with the demands of the Henderson-Hasselbalch equation. If the solution were not buffered, the change in pH on the addition of acid would correspond to the actual amount of added 114, in a buffered solution the change in pH is immeri-

 $\begin{array}{ccc} \operatorname{Hb} & + & \operatorname{O_2} & \rightleftarrows & \operatorname{HbO_2} \\ \operatorname{Reduced} & \operatorname{Oxygen} & \operatorname{Oxyhemoglobin} \\ \operatorname{hemoglobin} & \end{array}$

I rom this it can be seen that when the amount of free oxygen in the blood is lowered, as by diffusion into the tissues, the content of oxylicmoglobia will decrease and that of reduced hemoglobin will increase This occurs when arterial blood is changed into venous blood. When venous blood reaches the lungs, it is distributed in the many capillaries of the alveoli exposing a great surface to contact with the alveolar air The alveolar and is separated from the blood in the pulmonary capillaries by membrane at most 2 µ in thickness, this, in addition to the fact that crythrocytes with a surface area of some 80 square meters pass through the lungs each second, accounts for the rapid and efficient uptake of oxygen The oxygen tension of alveolar air (100 mm. Hg) is high enough to eause a considerable increase in the amount of dissolved oxygen in the bleod. This in turn in creases the amount of oxyhemoglohin at the expense of reduced hemoglohin at the expense of reduced hemoglohin globin, and when the aerated blood leaves the lungs as arterial blood practically all of the hemoglobin is normally in the form of oxyliemoglobin This relation between oxygen tension and the degree of oxygena tion of hemoglebin has been discussed in Chapter 22

Now in addition to their relation to oxygen transport, the preteins oxyhemoglebin and reduced hemoglebin aet as typical weak acids, being present in the blood partly in the unneutralized or free acid form (HHb HHbO3) and partly as salt ions (Hb-, HbO2) At the pH of the blood the reduced hemoglobin ion Hb- has a smaller base-binding capacity than has the oxyhemeglobin ion HbO2, ie, reduced hemoglobin behaves as 8 weaker acid than oxyhemoglobin When HbO2 is converted into Hb is the tissue capillaries by the loss of oxygen from the blood, the con comitant release of the extra base by the reduced hemoglobin would tend to make it slightly mere alkaline if it were not for the fact that carbonic acid is entering the blood at the same time. It has been shown that, under normal circumstances, about 50 per cent of the entering carbonic and is equivalent to this potential increase in alkalinity of the blood, 1 e, will be "neutralized" without any net pH change in the blood at all Since the carriage of this portion of the entering carbonic acid does not involve a change in pH of the blood it is known as the isohydric earnage aturally, the exact reverse of the above-described process occurs in the lung capil laries, where reduced hemoglobin is converted by oxygenation to oxygenation to oxygenation to hemoglobin This conversion increases the base-binding capacity of the hemoglobin withdrawing the base from BHCO₂ to form H₂CO₂, which be the standard of the s then decomposed to CO₂ and H₂O, the CO₂ diffuses out of the blood into the alveolar air of the lungs

It may be noted here that in addition to the carriage of a portion of the entering CO₂ by the isohydric reaction hemoglobin enters into carbon dioxide transport in at least two other recognized ways. One of these its act as a typical buffer in the mainter desembed in detail below. Inother is due to the fact that hemoglobin as a protein is capable of combining directly with carbon dioxide to form a carbamate the reaction involving

Chap. 24

the free amino groups of the hemoglobin molecule:

$$\begin{array}{c} H \\ HbNH_2 + CO_2 \rightleftharpoons HbN \\ \hline \\ COOH \\ \end{array} \rightleftharpoons HbN \\ + H^+ \\ \hline \\ COO^- \\ \end{array}$$

The formation of carbamate in this manner is a general property of proteins and amino acids and is not specific for hemoglohin, but, since hemo-

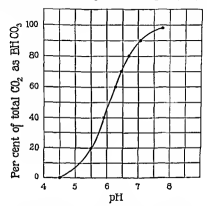


FIG 179. TITRATION CURVE FOR THE H₂CO₂ BHCO₂ SYSTEM. From Van Siyke Endocrinology and Metabolium, Vol 4, New York, Appleton, 1924.

globin is by far the most abundant protein in blood, a major portion of the CO₂ which is carried as carbamate is due to hemoglobin.

The existence of such a direct combination of hemoglobin and CO₂ was at one time disputed, but the work of Henriques, of Roughton, and of Stadie and O'Brien has established the significance of carbamate-bound CO₂ in respiratory gas exchange. While the actual amount of carbamate-bound CO₂ in the blood is small relative to the amount of bicarbonate (see Table on p. 687), the ease of formation and breakdown of this type of compound in response to changes in CO₂ tension is such that from 20 to 30 per cent of the extra CO₂ added to the blood in the tussues and released in the lungs may be transported in this manner.

Role of Buffers. As has been indicated, the isohydric reaction accounts for only part of the CO-carrying power of the blood. When more CO₂

cally much smaller, being equal to the change in the value of the logarithm of the ratio of buffer salt to buffer acid. It is for this reason that buffer solutions behave as they do in resisting gross changes in pH on the addition to the solution of acid or alkali, for it can be readily shown that an exactly analogous mechanism functions in the ease of added alkali

In a solution containing a number of different buffers, as is the case with the blood and tissues, the entering acid or alkali is buffered by all the buffer systems present, in proportion to their relative effectiveness at the given pH Thus in the buffer systems of the blood

B Protein вньо. B₂HPO₄ BHCO₁ BHbH₂CO₂ HHb BH.PO. H Protein HHbO,

the entrance of acid will eause a decrease in concentration of all the buffer salts (numerators) and an increase in concentration of all the buffer acids (denominators), accompanied by an equivalent change in pH as required

by the Henderson-Hasselbalch equation

It is well to note hero that physiological limitations influence consider ably the relative significance of the various buffers in the blood Thus the chief buffer for carbonic acid is hemoglobin, since carbonic acid is formed from CO2 and H2O only in the red cells where most of the buffering reac tions for carbonie acid take place (see below) On the other hand, the major buffering action for all acids other than carbonic acid is exerted by the plasma bicarbonate, protein, and phosphate buffer systems

It is significant that the pH of maximum efficiency of these buffers is

below the normal pH for blood, since as the blood pH falls, change of reaction is opposed with an efficiency which increases as the danger point is approached While the ratio BHCO, H2CO, at the pH of blood is about 20 1 and hence considerably removed from the ratio of maximum efficiency, the chief significance of this system rests in the fact that His Co. promptly desociates yielding CO2 which is expired, thus the actual effitiency of the CO2 system in neutrality regulation is greater than would appear to be the case from a consideration of the buffer theory alone

Role of the Red Cells. Serum separated from the red corpuscles ("separated scrum") has only a slight buffer effect and CO₂ carrying power as compared with scrum in contact with the cells ("true serum") Investigation has shown that this is because practically all of the reactions assor ciated with the transport of CO2 by the blood take place primarily within the red cell itself, the serum (or plasma) being involved in a secondar) marmer only The role of the red cell and of the plasma in the uptake of CO₂ by the blood is summarized in the accompanying diagram (Fig. 180)

As indicated in the diagram, CO2 produced metabolically in the tusues diffuses as such into the plasma. Here a small amount remains in physical solution, and some possibly reacts also with the plasma proteins (Pr 11) to form carbamate, the extra equivalent of acid thus formed being buffered by the plasma buffers (1-) in the usual way By far the greater portion of the entering CO₂ (upward of 90 per cent) does not remain in the plasm't, however, but diffuses rapidly through the red cell wall into the red cell realf

Here it comes under the influence of the enzyme carbonic anhydrase. This enzyme is present in abundance in the red cell (and in certain other specialized tissues, such as the pancreas and the gastric mucosa) but is absent from the blood plasma. Its function is to act as a catalyst in the reaction $CO_2 + H_2O \Rightarrow H_2CO_3$. By virtue of its presence within the red cell, much of the entering CO_2 is converted into H_2CO_3 , which immediately dissociates to give H^4 and HCO_1 ions. A portion of the entening CO_2 (about 20 per cent) escapes hydration, however, reacting rapidly with the hemoglobin present to form carbamate. The net effect of these two reactions, therefore, is the production within the cell of an excess of H^4 and HCO_7 ions, together with a small amount of carbamate ions. The

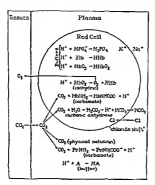


FIG. 180 PROCESSES INVOLVED IN THE UP-TIME OF CARBON DIOXIDE BY BLOOD

amounts of these ions thus produced are such as to greatly after the normal pH and osmotic equilibrium of the cell if they were not removed in some manner

The H+ ions are removed in two ways, by the isohydne reaction and by buffer action. It has already been pointed out that oxyhemoglobin (HbO₂) is a stronger and than reduced hemoglobin (Hb) at the pH of the blood. This means in effect that hemoglobin which has lost its oxygen has a lesser base-binding capacity at a given pH than has oxyhemoglobin. Thus the liberation of oxygen from the blood to the tissues creates a situation which brings about the release of base from the hemoglobin without involving any pH change at all. This process, under the conditions prevailing in normal blood, accounts for the disposal of over 50 per cent of the extra H+ ions produced within the rid call from the entering CO₂. The remaining H+ ions react with the various buffer sails ions present in the cell to form an equivalent amount of un-ionized buffer acids. Since

this typical buffer relation involves a change in the ratio of buffer salt to buffer acid, the plI of the cell contents will change as demanded by the Henderson-Hasselbalch equation, and it is this which accounts for the slight change in plI which follows the uptake of CO₂ by the cells

The fate of the extra IICO; ions must now be considered. If they remained within the cell, the increased osmotic pressure would cause the cell to take up water and swell to a size far beyond physiological limits. It has been found by chemical analysis of cells and plasma, that most of the extra IICO; ions diffuse out of the cell into the plasma, being replaced by an equivalent amount of Cl-ions which diffuse from the plasma into the cell. This process is called the chloride shift and it has been estimated that as much as 80 per cent or more of the critra brearbonate of venous plasma (as compared to arterial plasma) is due to diffusion from the red cells.

The chloride shift has been attributed by Van Slyke to the existence of a Donnan equilibrium (see p 12) across the red cell membrane It can be shown that, in the case of the red cell, it is n consequence of the Donnan theory that at osmotic equilibrium the ratio of bicarbonate-ion coaccurations in cell and plasma is proportional to the ratio of chloride-ion concentrations in cell and plasma, i.e.,

An increase of bicarbonate-ion concentration within the cell disturbs this equilibrium and it is partially restored by migration of bicarbonate ions from the cell to the plasma accompanied by an equivalent migration of helioride ions from the plasma to the cell 'A title same time, a certain amount of water diffuses from the plasma into the red cell, and this accounts for the slight increase in size of the cells of venous blood as compared to arternal blood.

The phenomena that have just been described take place during the uptake of CO₂ by the blood in the tissue capillaries to form venous blood. When the venous blood raches the lungs the entrance of oxygen reversed all of these processes. Bicarbonate in the cell is converted into carbonic acid which is dehydrated to CO₂ and diffuses out of the cells into the plasma, from whence it passes into the alveolar air of the lungs. Bicarbonate ions migrate from the plasma to the cell and undergo the same reactions, chloride ions returning from the cell to the plasma at the same time. The various other reactions are likewise reversed in an analogous manner. Thus the red cell is carried through the entire mechanism first in one direction and then in the other direction as it makes a complete circuit of the body.

The quantitative nature of the changes that normally take place in the

¹ The hemoglobia sons likewase enter into this equilibrium. For a detailed discussion of the Donana equilibrium as it is applied to the clioride shift, the reader is referred to Jeters and Van Styke Quantitative Clinical Chemistry Vol. I see also Hitchcock. Physical Chemistry for Students of Biology and Vedicine 3d cd.

blood in its transition from the arterial to the venous state is illustrated by the data of the following table, adapted from Stadie and O'Brien 2

-	Arterial Blood					Venous Blood					Difference									
	Ple	tema Cella		Il kele Blood		Plasma		Cells		li hole Blood		Plasma		Cells		Whols Blood				
	_		_																Per	en
Iematocrit	0	600	0	400			0	596	ļ٥	404			ı							
2 saturation per			96				•		74	.			1	- 1			1		ì	
H	7	45		12			۱,	13		11			ı	- 1			1		l	
CO mm ng	40		10		40	n	145		45		40	4	ı	- 1			1		ŀ	
Free CO: vol per			}			•	1	-	1	- 1		- 1		- 1			1		l	
cent	1	G	0	8	9	4	1	8	l٥	9	2	7	0	2 1	0	ì	0	3	8	0
Bound CO: vol per			Ì						П					1			1		1	
cent	34	1	11	8	45	8	36	3	13	1	49	4	2	2	1	3	3	5	92	0
B carbonate vol per			1				ļ		1	- 11				- 1			1		1	
cent	33	1	9	8	42	9	35	2	10	5	45	7	2	1	0	7	2	8	74	0
Carbamate vol per	١.		١.			_	Ι.		1	- 1		ш		- 1					} .	_
cent	1	0	2	0	3	0	1	1	2	6	3	7	0	1	0	6	0	7	18	0
Total CO: vol per cent	3.	_	10		48		38		14	. 1	52		2	. 10		4	١.	8	100	_

Study of this table shows the following

1 In its passage through the tissues of the hody, the arterial blood of the subject heing studied has pieked up about 38 ml of CO₂ per 100 ml of whole blood This is an increase of about 8 per cent in the total CO₂ content of the hlood With the R Q assumed to be 0.85, a normal value, this corresponds to the liberation of about 45 ml of O₂ from the hlood at the same time, which is responsible for the change in oxygen saturation from 96 per cent to 74 per cent

2 The increased CO₂ content of venous blood is accompanied by a rise in CO₂ tension of about 5 mm of Hg, and a fall in pH of about 0 01 to

0 02 pH umts

3 Of the total CO₂ of blood, almost 90 per cent is in the form of bearbonate, of which about three-fourths is found in the plasma and one-fourth in the cells. The remaining 10 per cent of the total CO₂ is about equally distributed between the forms of free CO₂ and carbamate. Plasma contains about twice as much free CO₂ and about half as much carbamate-bound CO₂ as is found in the cells

4 Of the 3 8 volumes per cent of CO₂ added to the blood in the change from arterial to venous blood, which are ultimately released in the lungs, about three-fourths appears as extra bicarbonate, about one-fifth as curbamate, and the remainder as an increase in the free CO₂ content. The extra free CO₂ and extra bicarbonate are both distributed between cells and plasma in about the same proportion as that already present, almost all of the extra carbamate is found in the cells.

^{*}J Biol Chem 117 439 (1957)

increased the hydron contentration of the blood or lowered its alkal reserve below the extreme normal limits." The conditions which fall into the different areas have been observed both clinically and experimentally

Disturbances of the Acid-base Balance. These are divided into two chineal types by Peters and Van Slyke, viz., (1) metabolic types, in which the primary disturbance is in the relation between alkali and acids other than H₂CO₂, and (2) respiratory types, in which the primary disturbance involves the CO₂ content of the blood in the table on p 691 the moe important conditions associated with disturbance of acid base balance are shown, together with their causes and the physiological mechanisms

brought into play for their compensation Kerosis Ketosis is the condition in which abnormal amounts of β -by droxybutyric acid, acctoacctic acid and acetone accumulate in the body fluids and may be readily detected in the unne. These three compounds are commonly known as the acctone bodies Of the three acctone bodies β-hydroxy butyric acid and acetoacetic acid are primary product. acctone being derived from the decarboxylation of acetoacetic acid Ther appear to have their origin principally from the metabolism of the fatts acids of fats, and to a lesser degree from certain amino acids resulting from protein cleavage This is discussed in detail in Chapter 33 They are apparently formed in the byer, and then further oxidized in the other tissues of the body where they are capable of furnishing a large amount of energy In diabetes mellitus, the body either does not possess the nor mal power of oxidizing these substances or else they are produced in excessive amounts At any rate, we find them in the blood and unae in abnormal quantity Likewise, in the absence of dietary carbobydrate and in other conditions of faulty carbohydrate metabolism, they are also in creased in amount in both blood and urine keto is as it appears in the human organism is not entirely the same as in other species, such as the rat, dog rabbit, goat, pig, and cow Rats are very resistant to the develop ment of ketosis

The significance of kctosis in connection with acid-base balance besides the fact that acetoacetic acid and β -hydroxybutyric acid exist in the blood and urine largely in the form of their alkali salts. Their production has therefore required an equivalent amount of body base for neutralization-Excessive amounts produce a severe demand on available base, and when they are excreted in the urine most of this base is lost to the organization of the produce of the product of the prod

The presence of acetone bodies in the urine in appreciable quantity and originally taken as an index of acidosis the severity of which was judged by the estimation of the amount of these substances present in the urine. That this is not a reliable index is shown by the occasional observation of a pronounced acidosis with no appreciable increase in urinary acetoric bodies. A high urinary ammonia coefficient (ammonia \(^1\) total \(^1\) national one looked upon as an indication of acidosis. However, this factor is not specific in diagnosis in spite of the fact that the majority of such cashow a high urinary ammonia value because certain dictetic changes may produce high urinary ammonia. Fatal aridosis has been observed in unitary ammonia fatal aridosis has been observed in the control of the

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2 Alkalı Reserve Direct Method

(a) CARBON DIOXIOE CAPACITY OF THE PLASMA (VAN SLYKE AND CULLEY) PRINCE PLE The plasma from oxalated blood is shaken in a separatory funnel filled with an ar imature whose carboo dioxide tension approximates that of normal arterial blood, by which treatment it combines with as much carboo dioxide as it is able to hold under oormal teosioo. A known quantity of the saturated plasma is then acidified within a suitable pipet and its carbon dioxide is liberated by the production of a partial vacuum. The liberated carbon dioxide is then placed under atmospheric pressure, its volume carefully measured, and the volume corresponding to 100 ml of plasma calculated.

Apparatus The apparatus used in the estimation of the carbon dioxide coatent of the plasma is illustrated in Figs. 182 and 183. It is notice of strong glass in order to stand the weight of mercury without danger of breaking and is held to a strong screw clamp the jaws of which are hood with thick pads of rubber. In order to prevent accidedtal slipping of the apparatus from the clamp an iron rod of 6 or 8 mm diameter should be so arranged as to project pader coef. If the very a could be

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It is necessary, of course, that both stopcocks should be properly greased and artight, and it is also essential that they (especially f) shall be held in place so that they cannot be forced out by pressure of the mercury. Rubber bands may be used for this purpose but clastic cords of fine wire spirals applied in the same manner as rubber bands, are stronger and more durable. Later models of this apparatus are equipped to overcome this difficulty.

Micr a determination has been finished the leveling bulb is lowered without opening the upper cock and most of fle mercury is withdrawn from the pipet through c. The water solution from d is readmitted and the leveling bulb being ruised to position 1, the water solution together with a little mercury, is forced out of the apparatus through a?

5 Most of the extra bicarbonate of venous blood is found in the plasma Now it has been shown that the pressige of blood through the capillanes is far too rapid to permit any appreciable hydration of CO, to H2CO (s necessary preliminary to the formation of bicarbonate) without the intervention of carbonic anhydrase, and this cury me is found only in the cell It follows, therefore, that most of the extra bicarbonate of venous plasma must have come from the cells. This is accomplished by virtue of the "chloride shift" mechanism, an equivalent amount of chloride leaving the plasma and entering the cells

Relative Importance of the Various CO2-Carriers of the Blood. \(\frac{1}{2} \) COrcarner of the blood has been broadly defined by Van Slyke as and substance present in the blood which increases the amount of CO: that may be taken up hy arterial blood without a change in pl1 beyond the normal difference between venous and arterial blood I rom this definition it is clear that the major CO-carriers of the blood include hemoglobin bicarbonate, the plasma proteins, and phosphates It is also clear that hy far the most important CO-carrier of normal blood is hemoglobia since 80 per ecut or more of the CO2 carriage by the blood is mediated directly or indirectly through the presence of hemoglobin. Thus this compound plays as important a part in neutrality regulation and the transport of CO2 as it plays in its more obvious function in O2 transport

Role of the Lungs. Due to the case of elimination of CO2 by respira tion, the regulation of the neutrality of the blood is largely controlled by the 112CO, BIICO, system From 20 to 40 hters of N/1 acid as CO, 18 lot each day by way of the lungs By means of variation in the rate and depth of respiration and the rate of blood flow through the lungs, opportunity afforded for very delicate adjustment of the hydrogen-ion concentration The nervous control of the respiratory mechanism resides in the respiratory tory center in the medulla Increases in CO2 tension or hydrogen-ion concentration call forth corresponding responses in the ventilation of the lungs, and since changes in either factor take place concurrently, it is difficult to distinguish the effects of one from the other. It is probable that an increase in hydrogen ions in the respiratory center itself, secondary to that of the blood, may he the activating factor Such increases may come ahout through diffusion of CO2 or H+ from the blood or hy acid production within the center The latter may explain the greater respiratory response which results from an insufficient oxygen supply to the center

The rate of elimination of waste products through the lungs and the kidneys is partly controlled by the rate of blood flow through those organs. The diminished blood pressure following excessive pulmonary ventilation has been shown to be due to the loss of CO₂ rather than to the resultant rise in pH per se There is, however, no constant relation between blood pressure and CO2 tension among different individuals

Role of the Kidneys. The role of the kidneys in neutrality regulation is concerned chiefly with the conservation of "fixed base" (sodium potassium) to the organism If a strong acid produced by metabolism within the tissues is neutralized by reaction with hiearbonate or basic phosphate to produce a salt, the hase component of the salt thus formed (eg, the Na of NaCl) is no longer a direct part of the alkali reserve of the body, and if present in excess will be excreted in the urine. If, however, the alkali reserve is low so that the organism needs to conserve fixed base, as when acid production exceeds the supply of available base, the kidney has the ability to excrete an acid urine (down to a pH as low as 4.8) as one means of conserving base. According to Pitts, this is probably accomplished by the reabsorption of base ions from the glomerular ultrafiltrate and their replacement by hydrogen ions secreted into the tubular lumen. In effect, the following reaction occurs:

$$Na_2HPO_4 + H^+ \rightarrow NaH_2PO_4 + Na^+$$
(from (from (to (to blood) kidney) urine) blood)

The increase in the relative amount of acid phosphate to basic phosphate in the urine accounts for the more acid pH of the urine. If the urine is titrated with alkah back to the pH of the blood (see "Thratable Acidity of Urine" in Chapter 31), a measure of the extent to which the above reaction has contributed to base conservation will be obtained.

A second and equally important mechanism whereby the kidneys conserve fixed base is by the synthesis of ammonia, probably from glutamine and amino acids. The ammonia is excreted in preference to fixed base, as follows:

$$NaCl + NH_4^+ \rightarrow NH_4Cl + Na^+$$
 (from (from (to (to blood) kldney) urine) blood

The extent of ammonia excretion is related to the need for base conservation, ordinarily being low or zero in alkalosis and bigh in acidosas. Thus by determining both the ammonia excretion and the titratable acidity of the urine, the extent of acid excretion by the body may be evaluated, and hence knowledge be gained concerning the state of acid-base balance.

It sometimes happens that there is an excess of base requiring exerction, as on diets high in alkaline ash or after administration of bicarbonate. In this event, extra base and bicarbonate will be found in the urine and the

pH may rise to as high as 80.

The PH and H₂CO₂:BHCO₂ Ratio. Inasmuch as the chief factor in the regulation of blood reaction is the H₂CO₂:BHCO₂ system, the relationship of variations in the latter to pH should yield valuable information concerning changes in the acid-base balance of the body. Applying the equation for the dissociation of a weak acid (see p. 682) to this case, the so-called Henderson-Hasselbalch equation is obtained.

$$pH = pK_1 + \log \frac{BHCO_1}{H_2CO_2}$$

The value of the constant pK_1 for blood plasma is 6.10. By measuring any two of the remaining variables, the third is of course determined. If we

³ Van Slyke, Phillips, Hamilton, Archibald, Futcher, and Hiller. J. Biol Chem. 150, 481 (1913).

plot changes in bicarbonate concentration [BHCO₃] as ordinates, against carbonic acid [H₂CO₃] as abscissas, a straight line will result for any given ratio of the two, and hence for any given pH. The slope of this line will increase or decrease with corresponding changes in the [BHCO₃] [H₂CO] ratio. The maximal as well as the normal ranges of these factors are depicted in Fig. 181, constructed by Vau Slyke. Since it was desired to use the customary form of CO₂ absorption curves, the coordinates are expressed in terms of total CO₂ values, [BHCO₃ + H₂CO₃], as ordinates and CO₃ tensions as abscissas It will be noted that time areas are set of by the three levels (high, normal, or low) of each of the two variables. The nine areas represent conditions of acid or alkali excess or deficit, which are

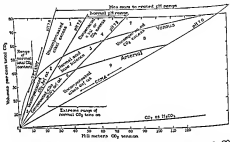


FIG. 181 NORMAL AND ABNORMAL VARIATIONS OF THE BHCO, H₂CO, CO: Tension and pH in Oxygenated Human Whole Blood Drawn from Resting Subjects at Sea Level. (Van Slyke)

The bicarbonate CO₂ at any point is obtained by subtracting from the total CO₂ the relatively small amount present as H₂CO₂ indicated by the slanting line near the bottom of the illustration

either compensated (pH normal) or uncompensated (pH above or below normal) That is, a condition of alkali deficit (low alkali reserve) may be either compensated by a corresponding diminution in carbonic and so that the [BHCO₃] [H₂CO₃] ratio remains normal (about 20 1), or it may be uncompensated by the failure to remove sufficient CO₂ in which case the pH falls and the condition of uncompensated alkali deficit results is therefore obvious that audosis cannot be regarded as merely a lower mig of the pH, since the aeidosis may accompany a deficit of CO₂ so marked as to result in a more alkaline reaction of the blood Vloreot of an abnormally acid reaction may occur even with an increase of alkali reserve provided CO₂ is present to excess In order therefore to combine the effects of both alkali reserve and reaction Van Slyke has broadly defined acidosis as a condition caused 'by the formation or absorption of acids at a rate exceeding that of their elimination [which] may be considered to have caused an abnormal state when it has either

DISTURBANCES OF ACID-BASE EQUILBRIUM OF BLOOD

Area (Fig IS1)	Acid-base Balance	Balance Conditions Associated Symptoms		Compensatory Mechanisms
I Uncompen- sated Alkalı Excess	[BHCO ₁] increased without propur- tionate rise in [H ₂ - CO ₂], therefore pH increased	Overdosage of Na- HCO: Excessive vanning (p; lotte ob- structura) or gastric lavage (loss of HCl). \(\lambda\)-ray ar radium treatment	1	Diminished respiration (rise in alveolar COi) to hold back CO-Diuresis and increased NaHCO excretion
2-3 Uncompen- sated CO ₂ Deficit	[H CO ₁] decreased without propor tionate fall in [BHCO ₄], there- fore pH increased	Hyperpnes, valua- tars or induced (axs gen want, eg, at high altitudes) Fever Hot baths		Retention of acid metabolites (low NH: and titratable acidity of urine) Excretion of Na HCO:
4 Compensated Alkalı of CO ₁ Excess	[BHCO ₄] (or [H-CO ₄]) increased but balanced by proportionate rise in [H ₁ CO ₄] (or [BHCO ₄]); there- fore pH normal	Allals excess NaHCO: therapy, with slow absorption CO: excess Retarded gas ex change (e.g., em physema) with CO: teasion chronically increased	Cyanosis due to deficient oxygen exchange	COs retention BHCOs retention
5 Normal	[BHCO _i] and [H _f - CO _i] normal at ordinary altitudes			
8 Compensated Alkalı or CO, Deficit	[BHCO ₃] (or [H ₇ CO ₅]) decreased but balanced by proportionate fall in [H ₇ CO ₃] (or [BHCO ₃]), there- fore pH normal	Alkals deficit Accelerated produc tion (c g , diabetes) or retarded elimina- tion (ca , nephritis) of nonvolatile acids Experimental acid in Diarrheal acidous of infancy (manamum) COs deficit Overventilation at high aligitudes (nx) gen want)	Нурегрпеа	Increased respira- tion (blowing off CO1) Accelerated NH1 formation and acid excretion Same as in Areas 2 and 3
7-8 Uncompen- sated CO: Excess	[H ₁ CO ₁] increased without prapor tionate rise in [BHCO ₃], there- fore pH decreased	Retarded respiration as in pneumons (physical abstruction) or morphine narcosis (desdening of respiratory center) Experimental rebreathing Cardiao decompensation	Dyspnea	Increased respira- tion Accelerated NHs formation and acid excretion Probable shift of acid from blood to tissue
9 Uncompen- sated Alkalı Deficit	[BHCO _i] decreased without propor- tionate fall in [H ₂ - CO _i], therefore pH decreased	Terminal stages of ne- phritic acidosis, and diabetic acidosis (compensated by in- sulin therapy) Deep ether aresthress Certain cardiae cases I clamposa	Dyspnes	Increased respiration Increased acid excretion and NIIs formation (except probably in nephritus)

increased the hydrion concentration of the blood or lowered its alkali reserve below the extreme normal limits." The conditions which fall into the different areas have been observed both clinically and experimentally

Disturbances of the Acid-base Balance. These are divided into two clinical types by Peters and Van Slyke, viz. (!) metabolic types, in which the primary disturbance is in the relation between alkali and acids ofter than II₂CO₂, and (2) respiratory types, in which the primary disturbance involves the CO₂ content of the blood. In the table on p. 691 the more important conditions associated with disturbance of acid-base balance are shown, together with their causes and the physiological mechanisms brought into play for their components.

brought into play for their compensation Kerosis Ketosis is the condition in which abnormal amounts of β -hy droxybutyrie acid, acetoacetic acid, and acetone accumulate in the body fluids and may be readily detected in the urine These three compounds are commonly known as the occione bodies. Of the three acctone bodies, 8 hydroxybutyric acid and acetoacetic acid are primary products acctone being derived from the decarboxylation of acetoacetic acid Ther appear to have their origin principally from the metabolism of the fatty acids of fats, and to a lesser degree from certain amino acids resulting from protein cleavage This is discussed in detail in Chapter 33 They are apparently formed in the liver, and then further oxidized in the other tissues of the body where they are capable of furnishing a large amount of energy In diabetes mellitus, the body either does not possess the nor mal power of oxidizing these substances or else they are produced in excessive amounts At any rate, we find them in the blood and urine in abnormal quantity Likewise, in the absence of dietary earbohydrate and in other conditions of faulty earbohydrate metabolism, they are also in ereased in amount in both blood and urine Ketosis as it appears in the human organism is not entirely the same as in other species, such as the rat, dog rabbit, goat, pig, and cow Rats are very resistant to the develop ment of ketosis

The significance of ketosis in connection with acid base balance hes in the fact that acetoacetic acid and β -hydroxybutyric acid exist in the blood and urine largely in the form of their alkali salts. Their production has therefore required an equivalent amount of body base for neutralization. Excessive amounts produce a severe demand on a valiable base, and when they are excreted in the urine most of this base is lost to the organization. The acidosis of diabetes mellitus may be largely due to ketosis. Acidosis occurs in many conditions however without a concomitant ketosis.

The presence of acetone bodies in the urine in appreciable quantity may originally taken as an index of acidosis the severity of which was judged by the estimation of the amount of these substances present in the urine. That this is not a reliable index is shown by the occasional observation of a pronounced acidosis with no appreciable increase in urinary according to the property amounts occificent (ammonia A total N) nast once looked upon as an indication of acidosis. However, this factor is specific in diagnosis in spite of the fact that the majority of such easies show a high urinary ammonia value because certain dietetic changes may produce high urinary ammonia. Fatal acidosis has been observed in uritima and in nutritional disorders of infants with no pronounced in

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Conditions of disturbed acid-base balance are best diagnosed and their course followed not by the determination of acetone hodies or ammonia in either urine or blood, but by the determination of certain other factors which are more or less typical of acidosis, such as the quantitative estimation of the alkali reserve of the blood and of its pH

METHODS

1. Simple Demonstration of the Presence of Carbon Dioxide in Expired Air. Into each of two small flasks or large test tubes introduce 25 ml. of a clear saturated solution of harum hydroxide. After an ordinary inspiration, expire through a hent glass tube or pipet dipped heneath the surface of the solution in the first flask. Repeat the experiment with the second flask, hut hold the breath as long as possible after the inspiration before breathing out through the tube Note the relative amounts of precipitate of harium carbonate formed. To another flask add 30 ml of water, 1 or 2 drops of barium hydroxide solution, and a few drops of 0.04 per cent phenol red Expire into this solution until a change takes place. What does this change indicate?

2. Alkalı Reserve Direct Method

(a) CARBON DIOXIDE CAPACITY OF THE PLASMA (VAN SLYKE AND CULLEW) PRINCIPLE The plasma from oxalated blood is shaken in a separatory funnel filled with an ar mixture whose carbon dioxide tension approximates that of normal attental blood, by which treatment it combines with as much carbon dioxide as it is able to hold under normal tension. A known quantity of the saturated plasma is then acidified within a suitable pipet, and its carbon dioxide is bhernted by the production of a partial vacuum. The liherated carbon dioxide is then placed under atmospheric pressure, its volume carefully measured, and the volume corresponding to 100 ml of plasma calculated.

Apparatus The apparatus used in the estimation of the carbon dioxido content of the plasma is illustrated in Figs. 182 and 183. It is made of strong glass in order to stand the weight of merenry without danger of breaking, and is held in a strong screw clamp the jaws of which are lined with thick pads of rubber. In order to prevent excidential slopping of the apparatus from the clamp, an iron rod of 6 or 8 mm diameter should be so arranged as to project under cock f between c and 6.

Three hooks or rings at the levels, 1, 2, and 3 serve to hold the leveling bulb at different stages of the analysis. The bulb is connected with the bottom of the apparatus by a heavy willed rubber tube.

It is necessary, of course, that both stopcocks should be properly greased and artight, and it is also essential that they (especially f) shall be beld in place so that they cannot be forced out by pressure of the mercury Rubber bands may be used for this purpose but clastic cords of fine wire spirals, applied in the same manner as rubber bands, are stronger and more durable Later models of this apparatus are equipped to overcome this difficulty.

After a determination has been finished, the leveling bulb is lowered without opening the upper cock, and most of the mercury is withdrawn from the pipet through a The water solution from d is readmitted and the leveling bulb being raised to position 1, the water solution, together with a little mercury, is forced out of the apparatus through a c.

⁴ It is well to have a funnel draining into a special vessel to catch the water residues and insecur; overflow from a \coma lerable amount of a creur; is it us regained if many analyses are run. It requires only straining through cloth or chain is skin to prepare it for

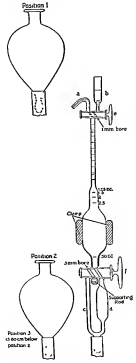


FIG 182 VAN SLIKE VOLUMPTRIC CARBON DIOXIDE APPARATUS.

Chap. 24

Procedure. Drawing the blood.⁵ About 6 or 7 ml. of venous blood are aspirated into a centrifuge tube (see Fig. 183) containing a little powdered potassium oxalate and some paraffin oil. The tube is subjected to a minimum of agitation after the blood is in it. The slight amount of agltation necessary to assure mixture with the oxalate is accomplished by stirring with the inleft tube, rather than by inverting or shaking. The tube and contents are then centrifused.

Saturation of Plasma with Corbon Dioxide. After centrifugling, about 3 ml. of the plasmar are transferred to a 300-ml. separatory funnel, arranged as in Fig. 184, and the air within the funnel is displaced by either alveolar air from the lungs of the operator or a 5.5 per cent carbon dioxide-air mixture from a tank. This latter procedure is preferred, since error due to incomplete filling of the separatory funnel with air containing carbon dioxide at the proper tension is less likely to occur. The gas must be passed through a wash bottle containing water before entering the funnel.

When alveolar air is used, the operator, with-



FIG 183 ASPIRATING TUBE USED TO COLLECT BLOOD

out inspiring more deeply than normal, expires as quickly and as completely as possible through the glass beads and separatory funnel. The stopper of the funnel should be inserted just before the

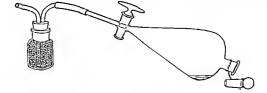


FIG. 184 SEPARATORY FUNNEL USED IN SATURATING BLOOD PLASMA WITH CARBON DIOXIDE. Courtesy, J. Bod Chem., 30, 289 (1917)

expiration is finished, so that there is no opportunity for air to be drawn back into the funnel. In order to saturate the plasma the separatory funnel

[•] For at least an hour before the blood is drawn the subject should avoid vigorous muscular exertion as this presumably because of the lactic and formed, lowers the bicarbonate of the blood.

^{*}II it is desired to keep the plasma for the estimation of carbon dioxide at a later time, it should be transferred to a parafilined tube, covered with a layer of parafilin oil, stoppered, and kept cold, under which conditions it is claimed that, if sterile it may be kept for over a week without alteration of its carbon dozinde expansity.

Is turned end over end for 2 minutes, the plasma heing distributed in a that layer as completely over the surface of the funnel's interior as is possible. After saturation is completed, the funnel is placed upright and allowed to stand for a few minutes until the fluid has drained from the walls and gathered in the contracted space at the bottom of the funnel.

Determination of Carbon Dioxide. The cup should be washed out with water and I ml. of distilled water run in, and together with the entire apparatus should be filled with mercury to the top of the capillary tube by placing the leveling bub of mercury in position 1. A sample of 1 ml. (of 0.5 ml. in case the amount of plasma available is very small) accurately pipeted, is allowed to run into the cup b in the apparatus represented in Fig. 182, the tip of the pipet remaining below the surface of the water as it is added, Add I drop of caprylic alcohol.?

With the mercury bulb at position 2 and the cock f in the position shown in the illustration, the plasma, water, and alcohol are admitted from the cup into the 50-ml. chamber, leaving just enough above the cock e to fill the capillary so that no air is introduced when the next solution is added in the case of whole blood, some corpuscies settle to the hottom of the cup. These are suspended in the water by stirring them up with the last 0.5 ml. of water after the first 0.5 ml. has passed into the chamber, Finally, 0.5 ml. of 5 per cent sulfurle acid is run in.

it is not necessary that exactly 1 ml. of wash water and 0.5 ml. of acid shall be taken, but the total volume of the water solution introduced must extend exactly to the 2.5-ml. mark on the apparatus, if the table on p. 700 ml. is

If the amount of plasma available is small, a little more than 0.5 ml, is saturated in a 50-ml, funnel, and exactly 0.5 ml, used for the estimation of carbon dioxide. In this case, the volume of distilled water and acid used wash the plasma into the apparatus is haired, so that the total volume of water solution introduced is only 1.25 ml, and in the calculation to describe of the carbon of the calculation to the carbon of the calculation to the carbon of the calculation to the carbon of the calculation to the carbon of the calculation to the carbon of the calculation to the carbon of the calculation to the carbon of the calculation to the carbon of the calculation to the carbon of the calculation to the carbon of the calculation to the carbon of the calculation to the carbon of the calculation to the carbon of the calculation to the calculation to the calculation to the calculation of the calculation to the cal

After the acid has been added, a drop of mercury is placed in b and allowed to run down the capillary as far as the cock in order to seal the latter. A copper wire may be used to expel a bubble of air which may be trapped in the capillary. Whatever excess of sulfuric acid remains in the cup is washed out with little water, using a medicine dropper.

The mercury bulb is now lowered and hung at position 3, and the mercury in the pipet is allowed to run down to the 50-ml. mark, producing a Torrice ilian vacuum in the apparatus. When the mercury (not the water) meniscrib has lailen to the 50-ml. mark, the lower cock is closed and the pipet is removed from the clamp. Equilibrium of the carbon dioxide between the 25 ml of water solution and the 47.5 ml. of free space in the apparatus is obtained either by mechanical shaking for 1 minute, or by turning the pipet upside down 15 or more times, thus thoroughly agitating the contents. The pipet is then replaced in the clamp.

Alter extraction of the gas the lower cock is opened, admitting the mercufinto the extraction chamber rapidly until the meniscus of the water solution

¹¹ is desirable to keep the amount of capelic alcohol small as larger amounts may appreciatly interase results. With plasma 002 not as sufficient to prevent foaming and is necessarist most consensually from a burst made by fusing a capillary stopcock to a plast graduated in 001-ml direction.

With whole blood in place of sulfuric and use lactic and made by diluting t volume of conventualed and top-such gravity 120) to 10 volumes with water

607

Chan 24

reaches the contracted upper portion of the chamber. At this moment the lower cock is partially closed and the remainder of the mercury is admitted nt n rate cufficiently retarded to prevent necillation of the water column in the calibrated portion of the apparatus when pressure equilibrium is reached The pressure is then adjusted by placing the mercury surface in the leveling bull above the moreury menicus in the chamber by a height equal to onethirteenth that of the water column, in order to balance the latter. After some practice controlled by a centimeter rule, one can estimate this level with the eye to within 2 mm of mercury, which is sufficiently accurate for most purposes. When the pressure has been adjusted, the lower cock of the apparatus is closed. The day volume may then be read at lelsure

When, as In most plasma analyses, the CO. Is not reabsorbed, no washing of the apparatus is necessary before using it for another determination, since the acid solution which were the walls of the chamber contains a negligible amount of CO.

CALCULATION For most chinical purposes, results within 1 or 2 volumes per cent of the true values for CO, capacity may be obtained by multiplying the observed volume (uncorrected) by 100 and subtracting 12. More accurate results are obtained by means of the table on p 700, in which the observed volume (corrected for pressure) is directly transposed into ml of earboa dioxide chemically bound by 100 ml of plasma. Tho value obtained from this table must be multiplied by 1 017 to correct for CO, re absorbed by the solution after release of pressure. The barometer reading and room temperature are taken at the time of the determination. For convenience in the calcu-

lation, values are given below for the ratio $\frac{B}{760}$ over the range usually encountered

In case the volume of plasma taken for estimation of earbon dioxide content was 0 5 ml , the observed volume of gas is multiplied by 2 before it is used to calculate the volume per cent of carbon dioxide bound

Interpretation. The carbon dioxide capacity of the plasma as determined by this method appears to indicate not only the alkaline reserve of the blood but also that of the entire hody The average normal value for man is 65 volumes per cent of carbon dioxide The table on p 701 shows the range of results obtained with normal and pathological plasma, as well as the relationship of the plasma bicarbonate to acid excretion, alkalı tolerance, and alveolar carbon dioxide tension

Barometer	B 760	Barometer	$\frac{B}{760}$
732	0 963	756	0 99a
734	0 966	758	0 997
736	0 968	760	1 000
738	0 971	762	1 003
740	0 974	761	1 005
742	0 976	766	1 008
744	0 979	768	1 011
716	0 982	770	1 013
748	0 984	772	1 016
750	0 987	774	1 018
752	0 989	776	1 021
751	0 992	778	1 021

(b) PLASMA BICARBONATE (TITRATION METHOD) VAN SLYKE! PRINCIPLE Plasma is treated with an excess of standard acid which is titrated back with standard all all to the original pH of the plasma as drawn

Procedure. Blood is drawn withnut stasis and without exposure to air into a glass syringe or tuhe coated with potassium uxalate and containing mineral oil (see Fig. 183). Then withnut exposure to air the blood is run into a tube under oil until the tube is completely filled. A one-hole rubber stopper is inserted into the tube, expelling through the hole the oil that remains over the blood. The hole is closed with a glass plug and the tube centrifuged. The plug is then taken out and as the stopper is removed from the tuhe, oil u allowed to run in through the hole in the stopper to cover the surface of the plasma, so that it is never exposed to air. The plasma is then transferred under oil to another tuhe. Simply covering the blood with oil is not sufficient to prevent loss of CO, during centrifuging.

A standard for the end point is prepared in accordance with Cullen's original directions for colorimetric hydrogen-ion determination. 20 ml. of neutral 0.9 per cent NaCl containing 7 drops of 0.03 per cent phenol red is placed in a round flask of about 100 ml. capacity and covered with a layer of mineral oil. I ml. of the plasma is then introduced under the oil and the

mixture is then stirred gently with a glass rod.

Another I-ml. sample of the plasma is transferred to a similar round flask, 5 ml. of 0.0f N HCl, which is made up in neutral 0.9 per cent NaCl, is added, and the CO, is removed by whirling the mixture vigorously about the flask for at least f minute with a rotary motion so that the solution is whirled in thin layer shout the inner wall. 10 ml, of 0.9 per cent NaCl and 7 drops of the 0.03 per cent phenol red solution are added and 0.01 N NaOH is run in from a huret, which permits readings to 0.01 mL, until the color matches that of the standard. As the end point is approached, sufficient 0.9 per cent NaCl B added to bring the volume to 20 ml.

The 0.01 N NaO11, like the 0.01 N 11Cl, is made up by diluting 1 volume of 0.f N solution to 10 volumes with neutral CO2-free 1 per cent NaCl. The use of saline solution instead of water has the advantage of preventing the forms

tion of a permanent cloudy precipitate of globulin.

A peculiar phenomenon occurs as the end point is approached. Each drop appears to change the color past the end point, but within a few seconds the color shifts back, and it is seen that at least another drop is needed. Const. quently, the final color comparison should not be made until at least seconds after the last drop of 0.01 N NaOlf has been added. When in double It is better to overrun the end point by a drop rather than stop short of it-

tracted from the ml. required to neutralize to the same indicator 5 ml of the 0 of N HCl used The number is approximately 5 but usually varies from it slightly because of difference in factors of acid and sikali and because of the calibration error of the 5-ml. pipet used in measuring the acid. The maximum accuracy is obtained by performing a preliminary titration on 5 ml of the acid plus 15 ml of distilled water, mark the same pipet, indicator, and end point as in the plasma titration. The titration reserves miles for a constraint of the constraint of th represents ml of 0.01 M NallCO, per ml of plasma and it is transformed into terms of molecular concentration of VallCO₃ by merely dividing by 100. If the NatlCO₃ molecular representation of VallCO₃ by merely dividing by 100. If the NatlCO₃ molecular concentration is multiplied by 2240 or the number of ml of 0.01 N HCl used in the titration by 22 t, the volume per cent of brearbonate CO; in the plasma is obtained and the results can thus be compared with those obtained by the CO₁method.

^{*} Van Slyke J Bod Chem 52, 495 (1922)

The standard 0.01 N NaOH must be protected from atmospheric CO, and kept in paraffined bottles to prevent solution of alkali from the wlass 10 The buret should be filled with fresh solution each day. The carbon te-free solution is made by dissolving the NaOH in an equal volume of H.O. On standing, the Na-CO, settles to the hottom 2.75 ml of the clear supernatural solution are diluted to 5 liters and standardized by titration with neutral red against 0.01 \ HCl. It is preferable to run the acid into the all all as the color change occurs without the time lag observed when all all is added to nord

Interpretation. The results agree closely with those of the CO. capacity method over the range of bicarbonate concentrations (0.03 to 0 01 M) ordinarily encountered in man, even in severe acidosis Below this range the titration continues to give accurate results, while the CO. capacity method gives somewhat higher values. For clinical and most experimental purposes, however, it appears that the two methods give results so nearly identical that they may be used interchangeably

3. Hydrogen-ion Concentration of Blood.

(a) COLORIVETRIC METHOD OF CULLEYII AS MODIFIED BY HAWKINS IS PRINCIPLE Blood is collected with precautions prainst loss of CO. The plasma is diluted with salino solution containing phenol red and the color obtained compared with that of standard phosphato mixtures of known hydrogen ion concentration

Procedure. Prenare two test tubes of the same diameter as have those used to contain the standards (see below), one containing 5 mil of 0.9 per cent sodium chioride solution and the other 5 mi of saline indicator solution,12 each covered with a i-cm layer of neutral mineral oil. Connect a Luer needle adapter, hy means of a short piece of rubber tubing, to a i-mi. Nohr pipet. Without stasis, insert the needle into the subject's vein, attach the adapter, and allow the blood to fill the pipet past the zero mark (A tourniquet may be used to ald in finding the vein, but the blood should flow into another tube a few seconds after the release of the tourniquet, so that stasis is absent when the adapter is inserted into the needle) Withdraw the needle from the vein and at once disconnect the rubber tubing from the pipet Wipe the tlp and Introduce into each tube, under the oil, 0.4 ml of the blood. Stir the contents carefully with a fine glass rod.

¹⁶ Testing Standard 0.01 N NaOH for Carbonate The solutions should be made up using only boiled water, be kept in paraffin lined bottles and be protected from atmospheric CO: by soda lime tubes They should be tested for carbonate as follows

To 5 ml of 0 01 N HCl in a 200-ml round fissk add from a freshly filled buret about 4 8 ml of the 0 01 N NaOH to be tested and 0.3 ml of neutral red solution. The mixture should be strongly acid to the indicator The solution is rotated for 1 minute in the flash to permit the e-cape of CO; and is then transferred to a 50-ml Erlenmeyer flask and titrated as in plasma analyses the total amount of 0.01 N NaOH required to give the end point being noted

A duplicate titration is performed in the same way except that there is no sgitation to remove carbon dioxide the 0.01 N HCl plus 20 ml of water being placed directly in the 50-ml Erlenmeyer flash, and the 0.01 N \a011 being added with a minimum of stirring

If there is no carbonate in the standard NaOH solution the two titrations give identical results. The difference should preferably not exceed 0 I ml and if it exceeds 0.2 ml the alkalı should not be used

¹¹ Cullen J Biol Chem 52, 501 (1922)
11 Hawkins J Biol Chem. 37, 493 (1923)

¹⁴ Prepare fresh for each determination Add 2 1 ml of 0 03 per cent phenol red solution to 100 ml of 0.9 per cent sodium chloride solution. Adjust to approximately pH 7 4 by starring with a fine glass rod dipped into 0 02 N alkali or acid, as the case may be

TABLE FOR CALCULATION OF CARBON DIOXIDE COMBINING POWER OF PLASMA*

Observed vol gas B	ol gas bonate by 100 ml			s brear Observed			MI of CO ₂ reduced to 0° 760 mm bound as bicar bonate by 100 ml of plasma			
× 760	15°	20°	25°	30°	^ 760	15°	20°	25°	30-	
0 20	9 1 10 1	9 9 10 9	11 7	11 8 12 6	0 60	47 7 48 7 49 7	48 1 49 0 50 0	48 5 49 4 50 4	48 6 49 5 50 4	
2 3 4	11 0 12 0 13 0	11 8 12 8 13 7	12 6 13 6 14 5	13 5 14 3 15 2	2 3 4	50 7 51 6	51 0 51 9 52 8	51 3 52 2 53 2	o1 4 o2 3 o3 2	
5 6 7	13 9 14 9 15 9	14 7 15 7 16 6	15 5 16 4 17 4	16 1 17 0 18 0	5 6 7	52 6 53 6 54 5	53 8 54 8	54 1 55 1 56 0	51 1 55 0	
8 9 0 30	16 8 17 8 18 8	17 6 18 5 19 5	18 3 19 2 20 2	18 9 19 8 20 8	8 9 0 70	55 5 56 5 57 4	55 7 56 7 57 6	57 9	56 S	
1 2 3	19 7 20 7 21 7	20 4 21 4 22 3	21 1 22 1 23 0	21 7 22 6 23 5	1 2 3	58 4 59 4 60 3	58 6 59 5 60 5	58 9 59 8 60 7 61 7	58 59 60 61	
4 5 6 7	22 6 23 0 24 6 25 5	25 2 26 2	24 0 24 9 25 8 26 8	24 5 25 4 26 3 27 3	4 5 6 7	61 3 62 3 63 2 64 2	61 4 62 4 63 3 64 3	62 6 63 6 64 5	62 63 64 65	
8 9 0 40	26 5 27 5 28 -	27 1 28 1	27 7 28 7	28 2 29 1 30 0	8 9 0 80	65 2 66 1 67 1	65 3 66 2 67 2	65 5 66 4 67 3	66 67	
1 2 3 4	29 30 31 32 33	3 30 9 3 31 9	31 5 32 4 3 33 4		1 2 3 4 5	68 1 69 0 70 0 71 0 71 9	68 1 69 1 70 0 71 0 72 0	68 3 69 2 70 2 71 1 72 1	68 69 69 70	
5 6 7 8 9 0 50	34 35	2 34 7 2 35 7 1 36 1 1 37	7 35 3 7 36 2 6 37 2 6 38 1	35 6 36 5 37 4 38 4	6 7 8 9	72 9 73 9 74 8 75 8 76 8	72 9 73 9 74 8 75 8	73 0 74 0 74 9 75 8 76 8	72 73 74 70 76	
1 2 3	40 41 42	0 40 0 41	4 40 9	9 41 2	1 2 3	77 8 78 7 79 7 80 7	78 8 79 6	79 6 80 6	77 78 79 80 81	
	42 43 7 44 8	9 43 9 44 9 45 8 46	3 43 4 3 44 3 45 2 46	8 43 9 7 44 9 7 45 6 46	5 9 6 8 7 8	81 6 82 6 83 6 84 5	81 5 82 5 83 4 84 4	84 3	S2 S2 S3	
0 6	0 4	3 8 47 7 7 48	1 48	5 48		85 8 86 3	86 2	86 2	80	

ids of columns represent in degrees centigrade the om ten perature at which the samples of columns represent in degrees centification and analyzed. It is assumed that the samples of plasma are saturated with alveolar carbon conde and analyzed. It is assumed that double and analyzed it is assumed that both operations are performed at the same temperature. The figures have been called a same temperature. perature. The figures have been so calculated that regardless of the room temperature which saturation and analysis are regional. which saturation and analysis are performed the table gives the volume (reduced to 700 mm) of carbon locate that the saturation and analysis are performed the table gives the volume (reduced to 100 mm) of carbon locate that the saturation and the saturation are saturated to 100 mm.) 700 mm) of carbon locade it at 100 ml of plasma are capable of binding when saturate at 20° with carbon dioxide at an 100 ml of plasma are capable of binding when saturated at 20° with carbon dioxide at anomaly. oce man ou carton norme that 100 ml of plasma are capable of bundars when saturate at 20° with carbon draude at approximately 41 mm tenson. If the figure in the table are multiplied by 0.94 they give within 1 or 2 per cent the carbon dounde bound at 37°

(a) Over 1 1 (b) Over 65

Good

(a) Below 20 mm (b) 27 per cent

direction§

(a) Over 100 (b) Over 6000 Over 6000

Below 30

10-30

Moderate to severe actevere acidosis Symptoms of acid intoxicadosis Symptoms may

be apparent

53-40

nounced symptoms

Laterne limits of bi-Mild acidosis, no pro-

Normal resting adult enrhonate reserve error in erther

Actual Sescree

Condition of Subject

Sodium Bicarbonale Required Вадаточкиг от тик Разки Вісавночате то Асір Ід**спеттон, Аблаі Тосьв**ансе, анр Абубобап Саввон Дісхірі. Тембіон* to I urn Urme Alkalıne (b) Approximate (a) G per kg || g for a 60-kg person May indicate | (a) 0 5-0 8 (a) 08-11 20-65 (b) 30-50 May indicate (a) 0-05 (b) 0-30 Corresponding Results of Indirect Tests for Acidosis dosis in its dosis than more act--נסט סננוספ Rehability in diabetes is present Carbon Dioxide of Alreolar Air absence Good (b) 68-47 per cent (a) 35-27 mm (b) 4 7-3 6 per cent (b) 3 6-2 7 per cent Approximate (a) Mm tension (a) 53-35 mm | Liable to con- | (a) 27-20 mm per cent m diabetes Rehability aderable 24-hr Excretion t of 0 I N acid + NII, Goods Good (b) Approximate (a) 27-65 (b) 1600-4000 (a) Ml per kg ml per 60 kg 1000-000 person (a) 0-27 (b) 0-1600 (a) 65-100 (b) 1000-600 ü Bicarbonale tol ver cent Brearbonate Reduced to 0°, 760 mm Plasma

rerity indicated by the corresponding plasma CO: figures in the first column

Ti e figures tabulated in this column also indicate the doses of bicarbonate necessary to restore the alkali reserve to normal from acidosis of the Measured either in 24 hour uring or in specimen from shorter period calculated to 24 hour basis Miter the irlanate administration likely to indicate more acidosis than is present * \an Siyko J Biol Chem 33, 271 (1018)

Centrifuge both tubes at about 1500 r.p.m. for 5 minutes. Compare with standardst in a comparator block, estimating between two standards, if necessary. The tubes are placed in the following relative positions:



The object of the saline plasma tube is to compensate the standard for the slight color and turbidity of the plasma (Walpole principle). The temperature of the saline indicator plasma is obtained, it is desirable to make determina-

¹⁴ Reagents and Apparatus Required Standards Sprensen's phosphate standards are prepared from Merck's special reagents in steps of 0.05 pH from about pH 72 to 71 ha M/15 phosphate solutions should be prepared from special reagent salts (Merck's are astifactory) by dissolving the following quantities in distilled water and making each solution up to one liter.

The proportions of acid and alkaline phosphates are given in the table. These mixtures may be kept for some weeks in pyrer glass in the refrigerator.

PROSPUATE MIXTURES (l'HENOL RED RANGE)

pH	M/15 NasHPO.	M/15 KH2PO4	μЦ	Ni/15 NazHPO.	M/15 KH ₂ PO ₄
	ml	ml		ml	ml
17 00	61 1	38 9	7 40	60 8	19 2
7 05	63 9	36 1	7 45	82.5	17 5
7 10	66 6	33 4	7 50	84 1	159
7 15	69 2	30 8	7 55	85 7	14 3
7 20	72 0	28 0	7 60	87 0	13 0
7 25	74 4	25 6	7 65	88 2	118
7 30	76 8	23 2	7 70	59 4	10 6
7 3.	78 9	21 1	7 75	90.5	95
		i	7 50	91.5	8.5

Color Mandarda These we prepared by adding 0.3 ml of 0.03 per cent phenol red (blends aulion) philaden) to 15 ml of each of the standard buffer solutions. The concentration of the required varies somewhat with different lots it is best to prepare a concentrated stock solution and determine by experiment the dultion required to gue axishactory depth of color over the desired pil range. The indicator solution must be neutral. After the addition of the color over the desired pil range. The indicator solution must be red.

The color standards must be renewed or checked against a fresh tube of die at least once every week as there is a slow fading of color.

Amandards The consumers.

Apparatus The comparator required as conveniently, made from a block 3 × 6 inches. The holes for the tubes are of 1 inch diameter. The shits for the light are best made by boring two 1; inch holes and going out the remaining wood with a classed. The tube

tinns at 20° C, by placing the plasma tubes, tagether with the necessary standard tubes, in a large heaker of water at 20°.

CALCULATION The pH of the blood (human) at 38° C is obtained from the following equation

$$pH_{16}$$
 = Colorimetric $pH_{t^0} + 0.01(t^0 - 20) - 0.23$

in which to represents the observed temperature. If the tubes are adjusted to 20° C, the middle factor naturally drops but

The empirical correction, -0.23 ("Cullen correction"), applied to adjust the colorimetric pH values to those determined electrometrically at 38°, compensates for the protein and salt errors in the colorimetric determination. The correction varies with the species of animal, but unfortunately is not as constant in each species as could be desired.

Hastings and Sendroy claim that the Cullen correction is unnecessary when the readings are made with the tubes at body temperature. Austin, Stadie, and Rohmson have shown, however, that under pathological conditions in man, and in the dog, there is considerable variation in the Cullen correction, and Hastings and Sendroy's procedure does not climinate it. For these cases, they state, corrections must be determined on each serum if colorimetric readings are to be relied upon

According to Cullen and Earle the colorimetric method gives results at 20° C 0.08 pH higher than the hydrogen electrode, and 0.14 pH higher than the quinhydrone electrode. The electrometric method using the glass electrode (sec (o) below) affords the most precise as well as most convenient method for measuring pH of the blood

Interpretation. The normal pH range of blood is between 7 30 and 7 50. The extreme limits of pathological variation which have been observed are 6 95 to 7 80. For the significance of abnormal pH values, see the table on p. 691.

(h) VETHOG OF SHOEK AND HASTINGS IN This is an adaptation of the preceding method employing a special pipet which not only permits the determination of cell volume as well as plasma p.H., but also makes possible the transfer of the diluted plasma to the manometric apparatus for CO₂ determination. Blood is collected by finger puncture under paraffin oil in the conneal receiver containing oxalato (Fig. 185). The Shock Hastings pipet is filled to the 0.1 ml math, followed by phenol red saline solution to the 2-ml mark. A control pipet is filled with 0.1 ml of blood diluted with indicator free saline solution. The tip of each pipet is covered with a bit of adhesive tape and a heavy rubber hand (Derhard Faber No. 84). The cell volume is read after centrifuging. The p.H is read in a special colorimeter block containing comparator.

must be of clear nonalkaline glass of uniform diameter. Tubes 20 min in diameter are

Light Either daylight or Daylite lamps are satisfactor;
Test for Ventrality The redistifled water is usually about pil 6.2 to 6.5 The easiest

test is that of using both phenol red and methyl red. The water should give no red color with either indicator. The syringe pipets and tubes should be rin-ed with redistilled water and dried by ringes.

the syringe pipets and tubes should be rined with redistinct water and dried syringes tubes and pipets washed and steribized in the usual manner employed in bacteriological laboratories are often dried from alkaline water

The saline solution must be adjusted to pil 7.4 as described above. The oxalate when dissolved in water to a 0.5 per cent solution should not be more alkaline than pil 7.2 to 7.4. The oil is tested by shaking with water contaming phenol red and methyl red. The water injust remain neutral.

¹¹ Shock and Hastings Proc Soc Exptl Biol Ved 26, 780 (1929) Also described in greater detail by Peters and Van Shjke Quantitative Clinical Chemistry Vol 2 p 804 Institutor The Williams & Wilkins Co. 1932.

tubes of the same internal diameter as the bulb of the pipets. The contents of the pipet may then be transferred to the Van Styke-Veill apparatus

(c) Electrometric Viltuon ¹⁸ The potentiometric method for the determination of pH has been applied to blood. The usual precautions must be observed in obtaining



Recenter for Blood

Shock-Hastings Pspet



Comparator Block

FIG 185 APPARATUS FOR SHOCK HASTINGS METHOD

the sample to prevent the loss of carbon dioxide, and similar care must likewise be taken during the actual measurement. To this end, glass-electrode vessels have been devised so that the sample of blood is maintained out of contact with air. Fig. 180



FIG. 186 GLASS-LLECTRODE AS SYMBLY FOR USE WITH BECKMAN PIL MLTER. Courtes) Finder Sea ent Se Company New

FIG 187 GLADS LLECTRODE FOR USE WITH THE COLEMAN ELECTROMETER.

Courtery A S Aloe Company St. Louis Mo

shows such a glass-electrode assembly for use with the Beckman pH meter—so designed that the sample of blood can be transferred with minimum gas exchange directly from it e hypodermic syringe to the electrode chamber. Anoth or model of glass electrode, for use with the Coleman Electrometer is shown in Fig. 187. In this assembly, the

^{*} Craig Lange Oberman and Carson Arch. Buchem and Buophys 38, 357 (19,2)

hypodermic needle is attached to the electrode chamber so that the sample of blood enters the chamber which is then immediately made part of the electrometric circuit. With the due care ordinarily exercised in the electrometric measurement of pH accurate values can be obtained on as little as 0 and of blood.

- 4 Acetone Bodies For methods of determining acetone acetoacetic and 8-hydroxybutyne acids in the blood see the References at the end of Chapter 23
- 5 Determination of Oxygen and Oxygen Capacity (or Hemoglobin) of Blood Volumetric Methods. It is possible to determine the oxygen of the blood using the same apparatus as that employed for the CO₂ estimation (see p. 693), suitable precautions being taken in collecting the blood for analysis. The oxygen content of blood represents the total volume of oxygen, both physically dissolved and combined with hemoglobin, present in 100 ml of blood in the condition in which it flows through the venso or arteries. For the determination of oxygen content it is therefore, necessary to collect the blood so as to avoid the oxygenating effect of air, i.e., under oil. An aspirating tube of the type illustrated in Fig. 183 is suitable for this nurrose

The exygen combined with hemoglobin in arterial or venous blood differs from the ovegen content in that a correction is made for free physically dissolved overen. The same precautions in the collection of the

sample must of course be taken

The oxygen capacity of blood represents the volume of ovygen required to combine with all of its hemoglobin (ox)hemoglobin and reduced hemoglobin). It therefore constitutes a measure of hemoglobin since it is established that 1 g of hemoglobin combines with 136 ml of ovygen or, in other words each ml of ovygen capacity represents 0.736 g of hemoglobin Blood for this determination may be collected without precautions to avoid contact with air, masmuch as aeration (ovygenation) is the first step in the procedure

The exigen unsaturation of venous or arterial blood is a measure of the absolute concentration of reduced hemoglobin and is obtained by subtracting from the exigen capacity (total hemoglobin), the exigen combined with bemoglobin as defined above. The distinction between this and the relative unsaturation, which is the per cent of total hemoglobin in the reduced state should be kept in mind. The use of the absolute rather than the relative expression is due to the fact that cyanosis is regarded as the result of an increased absolute concentration of the bluish reduced bemoglobin.

Determination of Oxygen Capacity Stir the special oxygen reagent (see p. 176) to obtain emulsification of the captylic alcobol Introduce 75 ml into cup b (Fig. 182), and deserate by shaking in the evacuated extraction cham ber Vix the blood thoroughly with a stirring rod and transfer to a 250 ml separatory funnel in which it is completely serated by rotating in a thin layer over the interior surface of the funnel (For oxygen determinations other than capacity, simply mix the blood under the oil and omit aera tion) Force 6 ml of the gas free reagent into the cup of the apparatus By means of a differential pipet (graduated to deliver between marks. Fig. 190), transfer exactly 2 ml of the blood directly into the extraction chamber by

keeping the tip of the pipet immersed to the bottom of the reagent in the cup and regulating the admission by the finger or stopcock on the pipet and cock e of the apparatus. Never allow more than a few ml, of blood to accumulate in the cup. An Ostwald transfer pipet may be used for this purpose, the final drop of blood being forced out of the immersed tip by expanding the air in the pipet with the heat of the hand, keeping the mouth of the pipet closed. Admit the remaining blood and all but I ml. of the reagent from cup b into the extraction chamber. With a medicine dropper discard the excess reagent and introduce a few drops of mercury as a scal. Evacuate and shake until the volume of extracted gas (O2 + CO2 + N2) is constant. This will be found to consume from 5 to 10 minutes and should be determined by measuring the gas at atmospheric pressure. Create a slight negative pressure within the pipet by bringing the level of solution to the 2-ml. graduation. Rinse cup b with distilled water and introduce 0.5 ml. of a 2 per cent NaOli solution previously deaerated. By means of the negative pressure within the pipet, admit this solution slowly, followed by a thin stream of mercury, which serves to break the column of NaOH which usually forms in the capillary of the pipet. Allow I minute for drainage and bring the gas (O: + N:) to atmospheric pressure as described in the determination of CO:, Read the volume of eas, the temperature, and the barometric pressure.

CALCULATIONS

$$V = \text{observed volume of gas } (O_2 + \mathbb{N}_2)$$

 $t = \text{temperature in }^{\circ} \mathbb{C}$
 $B = \text{barometric pressure in min. Hg}$

$$w = \text{tension of aqueous vapor (see p. 709)}$$

Volume per cent O₁ capacity =
$$\left(\frac{(B-w)}{760(1+0.00367t)} \times \frac{100V}{2}\right) - 21$$

Values for the temperature and pressure correction factor are given in the table on p 707. The following equation may also be used

Volume per cent
$$O_2$$
 espacity = $\left(\frac{179(B-w)V}{t+273}\right) - 21$

The value 2 1 in these equations corrects for ony gen and introgen play sically dissolved at atmosphene pressure 1 or oxygen content subtract 1 30 (per cent N₂) instead of 2 1 For O₂ combined with hemoglobin subtract 1 7 or 1 5 for arterial or venous blood respectively, which corrects for free O₂ and N₂ at arterial or venous tension O₃ unsaturation is obtained by subtracting O₃ combined with hemoglobin from O₄ capacity.

Hemoglobin (g. per 100 ml) = 0.736 × volume per cent O₂ capacity For colorimetric determination of bemoglobin, see p. 610

6. Determination of Carbon Monoxide in Blood—Volumetric Method of Van Siyke and Associates: Principle A more exact quantitative procedure for the estimation of carbon monoxide than the colorimetric method given in Chapter 22 is a volumetric method based on the experimenta of Van Siyke and his associates. The mixed gases are extracted from the sample of blood in a Van Siyke pipet (Fig. 182), the oxygen and carbon dioxide are absorbed and the carbon monoxide estimated in the readual gas either by absorption with Winkler's reagent or by correction for the nitrogen.

¹¹ Van Slyke and Salvesen J Bud Chem 40 103 (1919), Van Slyke and Stadie J Bud Chem 40, 1 (1921) Van Slyke and Neill J Bud Chem 61, 523 (1924) Adapted by Bernard L. Over

Time OF Figures FOR Circulation

Temperature	B-w 760(1 + 0 00367t) factor by which gas measured most at B mm is reduced	a cos	Arr† measured at room tem perature and pressure dis- solved by		$\begin{bmatrix} 1017f \left(1 + \frac{S}{50 - S} \alpha'_{CO1}\right) \\ factor by which the volume of CO obtained after 1 extraction is multiplied in order to obtain the volum of CO reduced to 0'' 760 mm contained in the solution analyzed$		
	to 0° 760 mm.*		2 o mi 11:0	5 tul H ₂ O	S = 2 5 ml	S = 50 ml	
• c			mi	ml			
15	$0.932 \times \frac{B}{760}$	1 075	0 0.2	0 10a	$1.002 \times \frac{B}{760}$	$1.061 \times \frac{B}{760}$	
16	$0.928 \times \frac{B}{760}$	1 043	0 051	0 101	0 995 × B	$1.053 \times \frac{B}{760}$	
17	0 924 × B 760	1 015	0 0a0	0 100	$0.989 \times \frac{B}{760}$	$1.046 \times \frac{B}{760}$	
18	$0.919 \times \frac{B}{760}$	0 989	0 049	0 098	$0.983 \times \frac{B}{760}$	$1.038 \times \frac{B}{760}$	
19	0 915 × $\frac{B}{760}$	0 966	0 048	0 096	$0.978 \times \frac{B}{760}$	$1.030 \times \frac{B}{760}$	
20	0 910 × B 760	0 942	0 047	0 095	$0.972 \times \frac{B}{760}$	$1.022 \times \frac{B}{760}$	
21	0 906 × B/760	0 919	0 046	0 093	$0.966 \times \frac{B}{760}$	$1.015 \times \frac{B}{760}$	
22	$0.901 \times \frac{B}{700}$	0 896	0 045	0 091	0 960 × B	$1.008 \times \frac{B}{760}$	
23	0 897 × $\frac{B}{760}$	0 873	0 045	0 090	0 954 × B	$1.001 \times \frac{B}{760}$	
24	0 892 × $\frac{B}{760}$	0 850	0 044	0 088	0 948 × #	0 993 × H	
25	$0.888 \times \frac{B}{760}$	0 828	0 043	0 0S6	$0.942 \times \frac{B}{760}$	0 986 × B	
26	$0.883 \times \frac{B}{700}$	0 808	0 042	0 084	$0.936 \times \frac{B}{760}$	0 978 × B	
27	0 878 × $\frac{B}{760}$	0 789	0 041	0 083	$0.931 \times \frac{B}{760}$	$0.971 \times \frac{B}{760}$	
28	0 873 × $\frac{B}{760}$	0 772	0 040	0 081	$0.924 \times \frac{B}{760}$	0 964 × B	
29	$0.868 \times \frac{B}{760}$	0 705	0 040	0 080	0 918 × B	0 957 × B	
30	0 863 × B	0 738	0 039	0 078	$0.912 \times \frac{B}{760}$	$0.950 \times \frac{B}{760}$	

* To calculate O; or hemoglobin when Or + N; volume is measured multiply gas volume by f to reduce to 0° 760 mm and by such factor as mecessary (100 when 1 ml of blood is used 50 when 2 ml are used) to bring results to volume per cent basis. Then for

1 7 val per cent N1 + dissolved O1

18 S

a O1 content subtract
b O1 bound by hemoglobus in venous blood sub-1 36 vol. per cent N.

^{1 5} vol per cent Nr + dissolved Or tract

O bound by hemoglobin in arterial blood subtract

Or bound by hemoglobia in blood saturated with air at 20° C subtract 1 vol per cent V₁ + dissolved O₂ = a 41d

Per cent of normal hemoglobin (Haldane scale) -

Grams of hemoglobin per 100 ml of blood = 0 736d Per cent of total hemoglobin saturated with $O_2 = \frac{100b}{100}$ or $\frac{100c}{100}$

Volumes per cent O: unsaturation = d - e nr d - b

I The d scole d arr is given as measured at room temperature. It is subtracted from it e air + COrolume measured after one extraction if plasma in aqueous carbonate solution, in order to obtain the COs, which is the numbipuled by 1 0.17 ($\frac{1}{10} - \frac{1}{10} - \frac{1}{10} - \frac{1}{10}$) an order to obtain the total volumes percent of CO₁ in the solution analyzed. When whele blood is analyzed, the air correction cannot be used because of the Op present and the CO1 must be determined by absorption with 1 and 1 to volume of gas absorbed in then multiplied by the above factor. The factor 1 0.17 being empirical, may vary abglith for inferent apparatus.

3 0 g.

8 0 g 4 0 ml

3 0 ml

Procedure. Blood should be collected under off in an oxalated tube. A special CO reagent" is required which differs from the O1 reagent in containing lactic acid as well as more potassium ferricyanide. The reagent (7.5 ml.) is made cas-free and the mixed cases are extracted from 2 ml. of blood exactly as described under the determination of oxygen capacity (p. 705). Longer shaking may be required to arrive at constant volume, since CO is not so readily dissociated from its combination with hemoglobin. After the extraction, a negative pressure is created in the extraction chamber by raising the solution level to a point 2 to 3 cm, below the 2.5-mi, graduation. About 0.5 ml, of mineral off is introduced into the cup and below this I mi. of alkaline pyrogaliate reagent.19 The reagent is admitted until its level reaches the capillary of the cup, and the column that forms in the graduated capillary is broken by means of a stream of mercury as described in the determination of oxygen capacity. A few minutes are allowed for drainage, the gas is restored to atmospheric pressure, and the volume read. The absorption of CO, and O: is repeated until the residual volume is constant. The CO may be estimated from this volume reading, the temperature, and barometric pressure.

If it is desired to determine the CO directly by absorption, proceed from this point by first drawing the mixed solutions in the pipet into reservoir d. This is to avoid mixing the alkaline pyrogaliate solution with Winkier's reagent to be used. Establish a slight negative pressure within the extraction chamber as described above. Rinse the delivery cup with distilled water and introduce into the pipet 0.5 ml. of Winkler's reagent. Absorption of CO takes place rapidly and the final gas volume should be read at once,

A method of magnifying small volumes of gas by reducing the pressure a definite amount below atmospheric is described by Van Siyke and Stadie. 11 Larger samples of blood may be used, the amounts of reagents being proportionately increased.

CALCULATIONS

or

V1 = Observed volume of gas from 2 mi of blood, after absorption by atkaline pyrogaliate solution

V2 = Observed volume after absorption by Winkler's reagent

B = Barometric pressure in mm of lig

w = Tension of squeous vapor in mm of Hg (see table on p 709)

t = Temperature in ° C

Volume per cent CO =
$$\left(\frac{(B-w)}{760(1+00367t)} \times \frac{100V_1}{2}\right) - 1.36$$

Volume per cent CO = $\left(\frac{(B-w)}{760(1+0)3670}\right)\frac{100(V_1-V_2)}{2}$

18 Special CO Reagent

Saponin Potassium ferricy anide Lactic acid c p Caprylic alcohol Distilled water to 1000 ml

" Alkaline Pyrogallate Reagent Prepare a solution of potassium hydroxide by dissolving 160 g. in 130 ml of water. In 200 ml of this solution dissolve 10 0 g. of pyrogallic acid-18 It ankler a Reagent Cuprous chloride 40 g ammonium chloride 50 g., distilled water to 150 ml

For use mix this solution with ammonium hydroxide (sp. gr. 0.9) in the proportion of 3.1.

" Van Slyke and Stadie J Bud Chem 49, 1 (1921)

Values for the temperature and pressure correction factor may be found in the table on p 707

VAPOR TENSION OF WATER

° C	mm Hg	°c	mm Hg
10	9 1	20	17 4
11	98	21	18 5
12	10 4	22	19 6
13	11 1	23	20 9
14	119	24	22 2
15	12 7	25	23 5
16	13 5	26	25 0
17	14 4	27	26 5
18	15 3	28	28 1
19	16 3	29	29 7
		30	31 5

MANOMETRIC METHODS OF ANALYSIS OF GASES IN BLOOD AND OTHER SOLUTIONS²²

Principle The manometric methods of gas analysis differ from the more commonly used volumetric methods in that the latter involve reading the volume with the gas under a definite (usually atmospheric) pressure whereas by manometric methods the pressure required to keep the gas at a fixed volume is observed. The advantages of this procedure are that a much lower degree of error is attinuable since in the former method the error in volume reading is many times greater than the error in reading of barometric pressure, calculation is greatly simplified as barometric pressure and corrections for vapor tension and capillary attraction of mercury do not enter in smaller quantities of material may be used and accuracy is attainable over a wide range of gas concentration.

Apparatus The apparatus coassis of a short pipet with the upper stem closed by a stopcock the lower connected with a glass tube. The latter descends then turns to connect with a leveling builb and a closed increury manometer. The pipet is call brated at two points to hold a ml of gas for pressure measurement and A ml of total volume respectively as shown in Fig. 188

For analysis the sample of blood or other solution is introduced into the chamber over mercury, together with the reagents to free the desired gises from combination A Torricellian vacuum is obtained as in the volumetric apparatus by lowering the leveling bulb, and the gases are extracted from solution by 2 or 3 minutes shaking The gas volume is then reduced to a m by admission of inercury and the reading p_1 is made on the manometer. The gases are either ejected or are absorbed by proper reagents and the reading p_1 is taken with the same gas volume. The partial pressure P of the gas at a ml volume is then $P = p_1 - p_2$ mm of mercury from which the gas volume at 0° 750 mm may be calculated. Methylene or ethylene glycol or glycerol may be used as a dehydrating agent to mositen the upper part of the manometer tube

³¹ Van Slyko and Neill J Biol Chem 81 523 (1924) Van Slyke J Biol Chem 73 121 (1927) In addition to the methods here to be described these papers give the details of the technique for the combined detarmination of all the gases in a sample of blood methods for micro-analyses of fractions of a ml of blood for determination of gases in liquid a saturated at high tensions for determination of dissolved gases in water and for to use of it apparatus in air and general gas analyses Principles and numerous applications of the manometric methods are completely described in Peters and Van 5\(\frac{1}{2}\) Quantitative Chinical Chemistry Vol 2 Baltimore The Williams & Williams Co. 19

The extraction chamber differs from that of the "volumetrie" apparatus in being calibrated at only three points, viz., 0.5, 2.0, and 50 ml. The mercury scal around the rubber joint at the bottom of the extraction chamber illustrated in Fig. 189 can be replaced by special heavy walled rubber tubbing A mechanical shaker is provided Air which diffuses through the rubber tubbing of the leveling bulb is expelled through

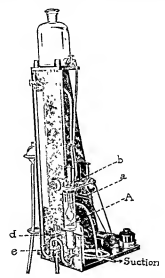


Fig. 188 Van Slike and Neill Portable Manometric Gas-Analysis Apparatus.

stopcock d The stopcock at the bottom of the manometer permits withdrawal of the mireury

The bottle on top of the frame holds dustified water. The lower bottle is to receive was the solutions drained out of the chamber after analyses. The most rapid and convenient way to transfer solution to the waste bottle is to force it up into the cup above the chamber and then draw it over into the bottle by suction. If suction is not available, a narrow rubber drain can be run from the curved outlet explicitly to the bottle.

Chap 24

necessarily maccurate modification of the manometric apparatus, see the original papers

Measuring Samples In order to attain the advantage of precision offered by the manometric apparatus accuracy in collecting and measuring samples is essential Blood should be either (1) drawn directly from the year without stasis, into the differential pipet from which it is to be delivered into the extraction chamber, or,

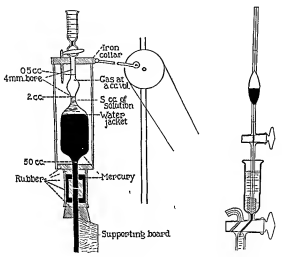


FIG 189 EXTRACTION CHAMBER

Fio 190 ATION OF DELIV ERY PIPET

better (2) collected under oil in an oxalated tube from which it can be drawn as needed The corpuscles should be uniformly distributed with a fine stirring rod before each sample is withdrawn A heavy walled Ostwald pipet graduated to deliver between two marks (differential' pipet) and preferably provided with a stopcock should be used (see Fig. 190)

Blood must be admitted at a rate slow enough to permit clear drainage of the pipet. Smooth delivery is usually obtained by using cock e or the cock of the pipet Amounts of solution of 2 ml or more may be run in under 1 ml of mercury in the cup the mercury then washing out the capillary For smaller amounts wash solution is necessury for complete transfer An ordinary transfer pipet may be used the final drop of blood being expelled by keeping the tip immersed closing the month of the pipet with

the index finger, and grasping the bulb with the warm palm of the other hand. The expanding air drives out the remaining blood

Cleaning the Apparatus. Preceding each analysis, the apparatus is cleaned by introducing 10 or 15 ml of approximately 001 \ lactic acid, shaking for 15 or 20 seconds in the exacuated chamber, and ejecting the extracted gas and solution Adherent solution introduces no error in the subsequent analysis.

FACTORS FOR CALCULATION OF VOLUMES PER CENT OF O1, CO AND N2 FROM PRESSURES IN 50-ML AFFARATES*

	Sample = 0 2 ml.	Sample = S =	- 1 ml - 3 5 ml	Sample = 2 ml $S = 7 ml$		
I emperature	S = 20 ml $a = 0.5 ml$ $t = 1.00$	a = 0.5 ml t = 1.00	a = 20 ml $a = 100$	a = 05 ml z = 100	a = 20 ml $a = 100$	
c						
15	0 312	0 0623	0 2493	0 0317	0 1251	
16	10	21	85	15	46	
17	09	19	78	14	42	
18	08	17	C8	12	37	
19	07	15	59	11	32	
20	07	13	50	09	28	
21	06	10	41	08	24	
22	0a	08	32	06	19	
23	03	06	23	05	15	
24	02	04	14	03	10	
25	01	02	06	02	06	
26	00	00	0 2398	01	02	
27	0 299	0 0598	, 90	0 0239	0 1198	
28	98	96	82	98	93	
29	97	93	74	96	89	
30	96	92	66	95	85	
31	90	90	58	94	81	
32	94	88	50	92	77	
33	93	86	42	91	73	
34	92	83	33	90	69	

[•] If cabbration of an apparatus shows a value of a significantly different from the 0.500 or 2.000 ml. in the column heading the factors in the column are corrected by multiplying them by $\frac{a}{0.000}$ or $\frac{a}{2.000}$.

Occasionally the apparatus shoul i be kept filled overnight with chromic-sulfung act that the admitted through cock b by regulation of cock e. The latter is kept closed but the scare open to allow the escape of gas

Testing for Leaks Test for leaks by introducing S ml, of water (Fig. 189) and extracting dissolved air for two moutes. Reduce the gas volume to a and read the pressure. If the temperature is constant, the pressure should not increase when this extraction is repeated.

 $S = 7.0 \, \text{m}$

Sample =

0 2 ml

FACTORS BY WHICH MILLIMETERS P_{CO} . THE MULTIPLIED TO GIVE VOLUMES PER CENT CO. IN SOLUTION ANALYZED-50-ML APPARATUS*

S = 20 ml

Sample = 1.0 ml

 $S = 3.5 \, \text{ml}$

ature	S=20ml					!	
	$a = 0.5 \mathrm{ml}$	a = 0.5 ml	a =20 ml	$a = 0.5 \text{ m}^{-1}$	a=20 m	a = 0 a ml	a=20 ml
	i = 1 037	i = 1 037	i=1 017	2=1037	i=1 017	i=1037	a=20 mi i=1 017
°C							
10	0 3454	0 0691	0 2710	0 0718	0 2818	0 0789	0 3097
11	37	87	0 2696	14	00	83	70
12	19	81	83	09	0 2783	76	-14
13	03	81	76	G _D	67	70	20
14	0 3386	77	56	01	50	64	0 2996
15	70	74	44	0 0697	35	58	74
16	54	71	31	93	19	52	50
17	38	68	18	89	04	46	28
18	22	64	06	86	0 2690	41	00
19	07	61	0 2594	82	75	36	0 2886
20	0 3292	58	83	78	62	31	66
21	78	56	72	75	1 8	26	48
22	63	53	60	71	34	21	28
23	48	50	48	68	20	16	08
24	34	-47	37	65	07	11	0 2790
25	20	44	26	01	0 2594	07	72
26	06	41	15	58	81	02	53
27	0 3193	39	05	55	69	0 0698	36
28 29	79	36	0 2494	ə 2	57	93	20
29	66	33	84	49	45	89	04
30	53	31	74	46	33	85	0 2688
31	-40	28	64	-43	22	S2	74
32	28	26	54	40	11	78	59
33	15	23	44	37	00	74	44
34	03	21	35	34	0 2489	71	30
To o the ml To c	n Slyke and S btain factor i of sample and alculate ral o	for a sample alyzed e.g.: f CO ₁ measi	other than for a 2 ml saured at 0° 7	I ml divid smple the fa 60 mm in	ie the above ctors are one the actual p	half of thes ortion of sol	e for 1 ml

lyzed use the above volume per cent factors for 1 ml samples divided by 100

Lubrication Cocks b and e must turn smoothly but not leak 1 thin layer of petrolatum is first applied followed by a rubber paste (made by dissolving 1 part of unvulcanized rubber in 5 parts of petrolatum with heat) using relatively less of the first coating in warm weather

Determination of Correction, c, for Manometer Depression Caused by Introduction of Absorbent Solution The introduction of absorbent solution causes a lowering of the mercury meniscus in the chamber and hence in the manometer, by

mereasing the volume of fluid between the mereury and the a mark at the moment of reading (Fig. 188). This necessitates for the p_1 reading a correction which is determined by blank analyses. The ship of the appiratism curves the area of the menuscus of the mereury in the chamber to v_1 recording to the values of S and a. The value of c for 1 ml of added solution may accordingly be from 1 to 1 mm, depending on the shape of the chamber and the volume of S.

Dilute absorbents (N NaOH) or hydrosulfite (20 per cent solution) have no significant effect on the vapor pressure in the chamber. Special precautions are required for alkaline pyrogallate solution (p. 708) because of the strong KOH

for alkaline pyrogallate solution (p. 708) because of the strong kOH.

When the final manometer reading is obtained after expulsion of the gases instead of after addition of an absorbing solution c, of course is zero.

CALCULATION The general equation for calculating total gas content of a solution from the volume of gas extracted in an evacuated chamber of definite volume as developed by Van Slyke and Stadie 2 has been subjected to certain modifications and made adaptable to manometric calculations.

Vapor pressure is eliminated as a factor since it is the same for both pressure readings and cancels out in the equation $P = p_1 - p_2$. An isoprection for reabsorption of extracted gas during the release of the vacuum is required for CO₃ just as in volumetre measurements and amounts to about 1014 For less soluble gases (O_3, N_1, CO) it is practically 1000 reabsorption being negligible A correction is also included for the effect of temperature on the specific gravity of increup:

The use of the rather involved final equation is expedited by the tables constructed by the authors which give directly the factors by which P read at any given term perature under the conditions ordinardly employed must be multiplied to obtain the volumes per cent of gas (see the tables on pp. 712 and 713). To express results as millimoles per liter eather use the table of millimole per liter factors given in the original papers or divide volumes per cent of gas by 2.24 or in the case of CO₂ by 2.246.

Volumes per cent gas =
$$P \times \text{vol}$$
 per cent factor m/s gas per liter = $\frac{P \times \text{vol}}{2.24}$

Determination of CO: in Blood or Plasma 24 The apparatus having been cleaned (see p 712), a drop of caprylic alcohol is drawn into the capillary above cock b, and 2 3 ml of CO, free water per ml of blood or plasma to be added are put into the cup Stopcock b is closed, with e open The blood or plasma is delivered beneath the layer of water in the cup from a pipet 28 described under "Measuring Samples ' After the delivery of the sample, the residue of blood in the cup is run into the chamber below, followed by the water layer Finally, 0 2 ml of CO, free 0 1 N lactic acld per ml of blood or plasma is added Stopcock b is then sealed with a drop of mercury The CO; is liberated by lowering the leveling bulb until the surface of the mer cury has fallen to the A mark, closing cock e, and shaking the mixture for 3 minutes The extracted gas is reduced to 2 ml (a), the admission of mercur) being regulated by stopcock e and the leveling hulb If the fluid meniscus passes this point, readjustment must be made by first bringing the mercury meniscus to the A mark and equilibrating for a minute Otherwise more re absorption of CO: will take place than is provided for in the calculation The adjustment being correct the manometer is tapped with the finger, and the height of the mercury column read (p, mm)

¹¹ J Biol Chem. 49 30 31 (1921) ¹² Shohl determines both CO₁ and pH on 0 1 to 0 2 ml of plasma. See J. Biol. Chem. 83, 759 (1929).

The variable amounts of O2 extracted from whole blood make it necessary to determine CO2 by absorption. This custom is also followed in plasma analyses, though it may be avoided by correcting for extracted air. The absorbent solution, 1 N NaOH25 is admitted under

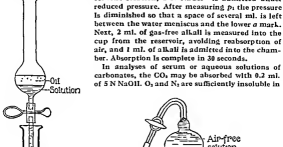


FIG 191. USE OF CAL-CIUM CHLORIDE TUBE RESERVOIR



FIG 192 BULB FOR PREPARING AND STORING GAS-FREE SOLUTIONS

this medium to make deaeration unnecessary. Only slight negative pressure is required to admit it. Air-free I N NaOH must be used for whole blood.

After absorption of CO₂ cock δ is seafed with mercury and the mentscus of the solution in the chamber is lowered a little below the a mark. Mercury is readmitted from cock e until the solution memseus is again on the a mark. Cock e is closed and the reading of the manometer is taken $(p_1 \text{ mm.})$. The CO₂ pressure $P_{co.}$ is

$$P_{CO_1} = p_1 - p_2 - c$$

where c is the correction discussed on p. 713. This correction is determined in a biank analysis in which S mi. of water, made alkaline with 2 or 3 drops of normal NaOil, are extracted in the apparatus, p_1 and p_2 being read before and after addition of the same amount of alkali as used in the determination.

$$c = p_1 - p_2$$

¹⁵ The alkali must be made gas-free and lept in properly protected reservoirs 25 to 30 m.l of 1 N NoOH are descrated in the extraction chamber and transferred under oil to a reservoir made from a calcium chloride tabe (Fig. 191). This will keep for a day. For larger amounts of solution, the apparatus isboan in Fig. 192 is suggested It consists of two leveling bulbs connected by a rubber tube 1 meter long. One of the bulbs, (diustrated) is provided with a disas expiliary and stopced. The bulb is half-dilled with absorbent solution and extracted by exacuation and shaking. Solution is delinered into the extraction chamber directly from the capillary of this reservoir.

To minimize error, all solutions must be measured very accurately, so that S varies within ±0 05 ml Also, sufficient time must elapse for temperature equilibrium to be established (For other sources of error and refinements of technique, see the original paper 19)

CALCULATION See p 714

Determination of Plasma CO; Capacity CO; capacity, as a clinical measure of acidois, may be determined by the volumetric method of Van Slyke and Cullen (p. 693). For the manometric determination, admit a drop of caprylic alcohol into the capillary of the cup, followed by 1.5 ml of 0.1 N lactic acid. One ml of plasma (previously saturated with alveolar air.) is introduced below the acid. The solutions are admitted to the extraction chamber, cock b scaled, the chamber exacuated and shaken 2 minutes. Then p. is read as described above with the gas volume at 2 ml. The gas is ejected, the ejected portion of the solution returned to the chamber, and the pressure reduced so that the gas space is again 2 ml. The value p, and the temperature are then read.

CALLULATION Pressure of extracted air and $CO_2 = p_1 - p_2$ The CO_2 capacity is obtained from this pressure by the use of the nomogram (Fig. 193)

Determination of Oxygen in Blood in Oxygen may be determined manametrically on 1 mi samples of blood with a variation within 0.2 volume per cent

The special O₁ reagent²⁴ is emuleified by rotating, and 7.5 ml are measured into the apparatue and deaerated by ehaking in vacuo for three minutes. During the shaking, the mercury chould be well in the neck below the bulb, to afford minimum exposure of surface, as emercury reacts with ferriegande. One mi of the well mitted blood to drawn into a pipet, preferably euch as described on p. 711.6 ml of the extracted reagent are forced into the cup, leaving 1.5 ml in the chamber. The blood is introduced under the reagent directly into the chamber as described under "Meaeuring Samples". The pipet is carefully withfavan and i ml of reagent is permitted to flow into the chamber, runsing through the blood in the capillary. The cock is sealed with a drop of mercury and the excess reagent discarded. The apparatus is evacuated and shaken for 3 mlnutes. I ml of air free I N haoll is placed in the cup and the CQ, absorbed by admixting 9.5 ml of the hydroxide into the chamber under diminished pressure as described in CQ, determination on whole blood.

The solution meniscus is brought to 2 ml (a) and p_1 (pressure of $0_1 + \gamma_2$) is read (For low 0) values it is preferable to read p_2 with a at 0.5 ml) λs in the volumetric method, 0_2 may now be determined either by making a correction for N_2 , or directly by absorption in the first case, after p_1 has been

³⁸ Austin J Biol Chem. 61 345 (1924) has described a molification of this procedure to be used in CO₁ analyses of serum obtained after ether anestles a.

to be used in COA analyses of serum obta ned after ether anestles a.

11 Sendroy J Bool Chem 91 307 (1931) describes a shorter method for hemoglobin
by manometric OA-capacity determination

Special Oxygen Reagent I otassium ferric; anide Saponin (Merck) Caprylic alcol 1 Water to

^{3 0} g. 3 0 g. 3 0 ml

Low results for on, sen may be accounted for 13 saponin of low 1 emolytic activity in which case the amount may be increased

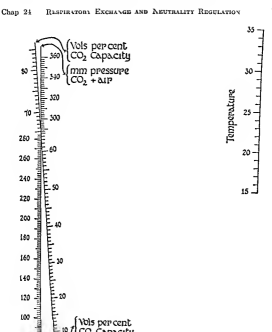


FIG 193 NOMOGRAM FOR CALCULATION OF PLASMA CO₂ CAPACITY
A strught line connecting the observed points on the pressure and tem
perature scales cuts the CO₂ capacity scale at the point indicating the
capacity

mm pressure

read, the gases are ejected from the chamber and p: is determined with the solution meniscus at the same a mark

The absorption method is resorted to for greater precision or when other gases (e.g., CO) are present After p, has been read, cock e is opened, with the leveling bulb in position to produce a gas space of 4 to 5 ml Cock e is then closed Next 15 ml of hydrogulfite solution. Is introduced into the cup

^{**} Grand up 100 g of sodium hydrosulfite and 10 g of sodium anthraquinone- β -sulfonate (Eastman Rodak Co) and keep in stoppered bottle To prepare absorbent solution stir

and by turning cock b admitted a drop at a time. As each drop trickles down, it absorbs O₂ and the mercury in the manometer falls. After a few drops have been admitted no further perceptible fall occurs. Cock e is then opened and the solution permitted to rise as near the stopcock as it will, with the leveling bulb in the lower ring. The remainder of 1 ml of hydrosulfite is then added, completing the absorption of traces of O₂. The gas is brought to the same a volume and p₁ is read on the manometer.

$$p_{0_1} = p_1 - p_2 - c_{0_1}$$

To check the completeness of Orabsorption and the absence of air leakage, expel the Nr and measure p. In the gas free apparatus. The Nr should approximate 12 volumes per cent in the Or content determination, or 14 volumes per cent in Oracapacity

An alternative absorbent for O₂ is the alkaline pyropaliate reagent (p. 708). This is not so clean or rapid as the hydrosulfite, but is more stable and its use is advised when O₃ determinations are done only occasionally. The gummy precipitate formed with the blood mixture necessitates mixing the solutions in the bulb of the chamber, well below the a mark, resulting in a greater negative pressure for the admission of the absorbent. One mill of alkaline pyropalitate is admitted dropwise over 4 or 5 minutes. The pressure is then allowed to rise to atmospheric so that the upper tube is rinsed by the solution, thus avoiding a reduction of vapor tension due to the strong alkali. Then p₂ is read. The apparatus is cleaned with water after using this absorbent.

The c correction, when oxygen is determined by absorption, is due to two factors, the lowering of the mercury meniscus by the absorbent and the race of O_1 left in the reagent after one extraction T0 determine c, describe 7.5 ml of the special O_1 reagent for 3 minutes, expel 5 ml, leaving 2.5 in the chamber. Shake again for 3 minutes and then admit 1.5 ml of air-free 1.8 NaOll in the manner described for CO_2 . Read p_1 , with a at 0.5 and 2.9 ml. Then run in the absorbent as in the analysis and obtain p_2 at both points. Use CO_2 recresponding to a in the analysis.

$$c_{0_1} = p_1 - p_2$$

Eject residual No and read po

$$c_{h_1} = p_1 - p_1$$

If O_1 is determined indirectly by correcting for \searrow_1 instead of by absorption, cls obtained in the same way, except that p_1 is measured after ejection of the $O_1+\searrow_1$, without preliminary absorption

CALCILATIONS. The estimated corrections for dissolved O₁ and N₂ of blood in calculation of total or combined oxygen or oxygen capacity given in the accompanying table, may vary somewhat with the cell content

The corrections are subtracted from the volume per cent of gas as determined by the use of the table on p 712

¹⁰ g. of this matture in 50 ml of 1 \times KOH and quickly filter it rough cotton. Describe solut in and transfer to container under oil as deverted in it is footnote for p 710. One drop of 10 per cent i cCh still further accelerates the absorption activity. The solubility of \(\times_1\) in this medium is negligible.

Chap 24

ESTIMATED CORRECTIONS FOR DISSOLVED O2 AND N2 IN BLOOD

Blood	Determined	Sought	Correction to Subtract		
			Vol per cent	If per l	
Venous	Total Os	Combined O2	0 1 (01)	0 04 (O)	
Arterial	Total O1	Combined O ₁	0 2 (O ₂)	0 09 (O1)	
Saturated with air at 20° 760 mm	Total O:	Combined O2 (O2 ca	05(0)	0 22 (O ₂)	
Venous	Total O: + N:	Combined O2	13 (O ₃ + N ₂)	0 a7 (O: + N:	
Artenal	Total O1 + N1	Combined O ₂	1 o (O ₂ + N•)	0 62 (02 + 32	
Saturated with air at 20° 760 mm	Total O2 + N2	Combined O ₂ (O ₂ ca pacity)	1 9 (O ₂ + N ₂)	0 85 (O1 + N1)	
Venous	Total O2 + N2 or CO + O2 + N2	Total Or or CO + Or	1 2 (N ₂)	0 53 (71)	
Artenal	Total O1 + N1 or CO + O1 + N1	Total O1 or CO + O1	1 2 (N ₂)	0 53 (Nz)	

Determination of Carbon Monoxide Hemoglobin, of Methemoglobin, and of Hemoglobin by the Carbon Monoxide Capacity Method. Principle Carbon monoxide is liberated from combination with hemoglobin by treatment with an acid ferrey unide solution CO₂ and O₃ are absorbed with alkaline pyrogallol solution and CO determined by correcting for N₂ in the residual gas

For determination of bemoglobin the blood is saturated with CO and the CO capacity determined Volumes of CO absorbed are identical with those for O₁. This gives active bemoglobin

CO without the sid of hydrosulfite does not change metheinoglobin to carboxy hemoglobin. The difference between CO capacity with and without hydrosulfite treatment represents methemoglobin. Hemochromogen behaves like methemoglobin hut is rarely present **

Determination of Hemoglobin (Active) by CO Capacity Draw 1 drop of caprylic alcobol into the capillary beneath the cup of the manometric apparatus Measure 4.75 ml of water into the cup. With stopcock pipet provided with rubber tip (see p. 711), run 2 ml of blood directly into the chamber, followed by the 4.75 ml of water. Place 1 to 2 ml of mercury in the cup above the chamber. Fill the outlet capillary of a Hempel gas pipet containing CO (see Fig. 194) with mercury. Fit the tip of the pipet into the bottom of the cup as shown in the illustration. Turn the two cocks shown in the illustration so that CO gas³¹ can flow from the pipet into the chamber. The flow is regu-

²⁰ Van Slyke and Hiller J Biol Chem 78 S07 (1928) 84, 20o (1929) Methods for smaller amounts of blood are also given For a somewhat more accurate procedure for the determination of carbon monoxide in blood see Sendro, and Liu J Biol Chem 89, 133 (1930)

³¹ Conant Scott and Douglass J Bul Chem 76 223 (1928) give a method suitable in the presence of hemochromogen

³¹ The bulbs are of about 30-ml capacity each. The capillary is of 1 mm hore. When he pipet is not in use a little mercury is let into the capillary leading to the CO bulb to seal the three way cock and prevent leakage around it. This drop of mercury in the capillary to the right of the cock is shown in the illustration. This project may also be used to store are free solutions. In this case the solution replaces the CO gas shown above, and mercury is used where water is indicated in the above figure. Supplied by the inviters of the Van Shk-Nell apparatius.

^{**} CO Gas. Connect the lower openings of two 5-liter aspirator bottles with rubber tubing at least 16 nm wide Fill one (1) completely with water 1 is 4-lo fitted with a thistle tube with stopeock and a side tube with stopeock connecting with a large test tube carrying a 4-dity thistle tube containing incremy possing through the stopper only and a

lated by the cock leading to the mercury leveling bulb of the manometric apparatus. With the leveling bulb in mid-position (a little below the bottom of the chamber), open this cock slowly, withdrawing mercury from the chamher until CO enters to the 2-ml, mark. Close the cock of the chamber, seal with a drop of mercury, and evacuate. With the mercury meniscus at the

TIO 194 HEMPLL PIPLT PRO-VIDED WITH THREE-WAY STOPCOCK15 (VAN SLYKE AND HILLER)

50-mi mark, shake until equilibrium is reached (i minute or longer as determined in blank below). Eject the mixture of gases from the chamber.

Evacuate until blood solution is in lower fourth of chamber. Put about 1 ml. of mercury and 2 to 3 ml. of water in the cup. Through the mercury scal introduce 0.25 ml. of acid ferricyanide24 solution, using a rubber-tipped buret made by fusing a stopcock onto a pipet graduated in 9.91-ml, divisions. Before the tip of the buret is inserted into the mercury, move It through the water layer to disiodge ferricyanide crystals or air bubbles. Fill the capillary and bore of the cock with mercury but run none into the chamber. Evacuate, lowering the mercury to the 50-ml. mark. Shake slowly for about 5 seconds, and then vigorously for 3 minutes. Admit mercury until gas space is reduced to 5 to 6 ml. Measure 2 ml. of air-free

I N NaOIIst into the cup, letting it run in slowly with the tlp against the bottom of the

cup. Allow I mi. of the NaOII to flow slowly Into the chamber, Absorption of CO, is complete in less than 1 minute. Bring the volume of gas to 2 ml Record the manometer reading as pr. Eject the gas. Seal the stopcock with a drop of mercury. Lower the fluid meniscus to 2 ml. Read the manometer again (p.).

CALCULATION. The hemoglobin content of the blood in terms of CO- or Opcombining power is calculated by the equation

CO or
$$O_2$$
 capacity = $(p_1 - p_2 - c)f$

where f is a factor obtained from the table on p 712 and c is determined by a black analysis in which the procedure described above is repeated in every detail except that 2 ml of water are substituted for the 2 ml of blood. The correction is calculated as $c = p_1 - p_2$ The value of c_i once accurately determined, can be used as a constant since it is small and insensitive to ordinary changes in laboratory conditions

Determination of Total Hemoglobin and Methemoglobin. Wash apparatus with three successive portlons of 10 to 15 ml of water, to the first of which

second thisti, tube with stopcock reaching to the bottom of the test tube. Run into the test tube 3 ml of anhydrous formic acid and then allowly concentrated HiSO, gently warming the mixture with a microburner When about 300 ml of CO and air have collected in A drive this out by opening the thistle tube and lifting B Continue generation until the CO from all the forms: and is collected in A. Clamp the tube between the two bottles with a screw clamp and detach the test tube. (arry out the entire procedure in a hood or in a free draft of air Smaller vessels may be used if less CO is desired

" lend berricyaride Solution To 92 volumes of stock solution, containing 32 g of KiFel(N)s per 100 ml add 8 volumes of concentrate lastic and of sp. gr. 1.2. It may be used for more than 2 months. 1 N VaOII Rendered are free as described up 715 a little hydrosulfite solution is added. This is done by evacuating to the 50-ml mark running in the water, shaking for 15 to 20 seconds and electing the colution

Draw 2 drops of caprylic alcohol into the capillary beneath the cup. Measure into the cun 4.3 ml. of water. Using the stongack ninet with rubber tin run 2 ml, of blood directly into the chamber, followed by a few drops of the water in the cun to wash blood through the capillary. From a microburet (see above) run 0.4 ml of the ammoniacal sodium hydrosulfites solution into the chamber, followed by the water remaining in the cup. Put 1 to 2 ml. of mercury in the cup. Lower the mercury in the chamber to the 50-ml. mark Run in CO from the Hempel pipet until the pressure on the manameter rises about 150 mm, the mercury remaining at the 50-ml, mark, if 100 mm of CO are run in and then mercury admitted from the leveling bulb to raise the level to the 50-ml, mark, the proper pressure will usually be attained Shake for 136 minutes, Eject gases from the chamber, Determine CO exactly as in the method above for active hemoglobin, except that 0.30 ml. instead of 0 25 ml, of the ferricyanide solution is run in. Determine correction c by cumping a blank test on 2 ml. of water (see above). This c will be somewhat greater than in the method for active hemoglobin. The calculation is otherwise the same and gives total hemoglobin.

Methemoglobin = total hemoglobin - active hemoglobin

Manoinetric Determination of Other Substances, See footnote 22, p. 709 Other manometric methods are those for urea using the ureasess or hypobromites, procedures oxalic acid or calcium. ** total reducing substances ** and fermentable sugars to in blood and urine, lactic acid in blood, 41 potassium in serum, 48 gas in fermentation,48 primary amino nitrogen,44 cysteine and cystine,48 gas mixtures,46 carbon in organic substances,47 chloride in serum and unne 48 phosphorus in organic compounds.49 carboxyl groups in amino acids,50 inorganic sulfate,51 magnesium,52 amino acids in blood, \$2 amino acids in urine. \$4 premase, \$5 total carbon and its radioactivity \$4 For some of these methods, later references will be found in Chapters 23 and 31

²⁶ Ammoniacal Sedium Hydrosulfile Pour 50 ml of chluted (1 50) ammonia solution upon 20 g of pulverized Na:S:O: contained in a 100-ml beaker Cover at once with a layer of paraffin oil Dissolve by stirring with a rod for a few seconds

³⁴ Van Slyke J Biol Chem , 73, 695 (1927) 17 Van Slyke J Biol Chem. \$3, 449 (1929)

Van Slyke J Biol Chem, 33, 445 (1929)
 Van Slyke and Sendroy J Biol Chem 34, 217 (1929)
 Van Slyke and Hawkins J Biol Chem 79, 739 (1928)
 Van Slyke and Ilawkins J Biol Chem, 33, 51 (1929)

⁴¹ Hastings and Avery J Bul Chem 94, 273 (1931)
42 Kramer and Gittleman Proc Soc Expd Bul Med, 24, 241 (1926)

^{**}Raymond J Biol Chem 83, 611 (1929)

**Van Slyke J Biol Chem 83, 425 (1929)

**Baernstein J Biol Chem 89 125 (1930)

**Van Slyke and Hanke J Biol Chem 95, 569 587, 599 (1932)

⁴⁷ Van Slyke Page Irvine, and Kirk J Biol Chem , 100, zeni (1933)

^{**} Sendroy J Biol Chem 109, lxxx (1935)
** Kirk J Biol Chem 106 191 (1934)

Van Slyke, Dillon MacCayden and Hamilton J Biol Chem 141, 627, 671 (1941)
 Hosgland J Biol Chem 136, 543 (1940)
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 Van Siyke, MacFayden, and Hamilton J Buol Chem. 150, 251 (1943) 35 Van Slyke and Archibald J Biol Chem , 165, 293 (1946)

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25

Energy Metabolism

Historical. Combustion in living beings was first described by Lavoisier (1743-1794) This brilliant chemist discovered the true nature of oxygen, and how its union in the body with carbon and hydrogen resulted in the formation of carbon diovide and water and in the production of beat In experiments on man he determined that oxidation was increased by food, by a cold environment, and by muscular work. He knew the main constituents of the atmosphere and that nitrogen played no active part in animal respiration. Lavoisier was under the false impression that oxygen decomposed some fluid in the lung causing the liberation of by drogen and carbon with the subsequent oxidation of these elements and their excretion as carbon dioxide and water in the expired air Scientists of a later date abandoned the theory of ovidation in the lungs, and favored the blood as the site of these chemical changes. This belief was strength ened by the discovery of gases in the blood by Magnus in 1837 Later it became known that, for the most part, oxidation occurs in the tissues and that the blood simply carnes the gases to and from the lungs

In 1842 Liebig announced that the substances burned in the body were carbobydrate, fat, and protein, and suggested that the unmay introgen would serve as an index of the extent of protein destruction. Ten years later Bidder and Schmidt described protein metabolism, and shortly afterwards Carl Voit measured protein waste by determining the amount of introgen in the excreta. You made the important discovery that muscular exercise did not increase protein metabolism. During many years of careful work, Rubner, a pupil of Voit, determined the fuel values of the foodstuffs and by means of a respiration calorimeter showed that the heat production as calculated from the respiratory exchange was the same as that obtained by the direct measurement of the heat given off by the body. In 1883, Rubner demonstrated the relationship between the surface area of the body and the heat production, thus providing a basis for comparison of the metabolism of different individuals.

In 1915 DuBois devised the most satisfactory method for estimating the surface area of the body, and later published normal standards of heat production for males and females, the accuracy of which has been fairly well established

In America, Atwater, Armsby, Benediet, Lusk, DuBois, and their many associates have added valuable contributions to the knowledge of normal metabolism. Benedict and those associated with him in the Carnegie Nutrition Laboratory have in addition perfected various forms of respira-

723

tion apparatus, and they are chiefly responsible for the extensive use of calorimetry in the clinic

Physical and Chemical Bases of Animal Calorimetry.

Gases are compressible fluids. The molecules of which they are composed are in constant motion and tend to disperse. The gas therefore expands and fills all parts of the containing vessel and exerts pressure upon the enclosing walls. Heat increases the tendency to expand and so increases the pressure unless the container permits the gas to assume a larger volume. If the temperature remains constant and pressure is applied to the container, the molecules are forced closer together and the volume is decreased. Therefore, when we speak of a volume of gas, we must also state the temperature and pressure under which it was measured. Regardless of the experimental conditions of measurement, gas volumes are always recorded as the volume at standard temperature (0° Centagrade or 273° Absolute) and standard pressure (750 mm of mercury at 0° C !). This is sometimes referred to as standard conditions or as N T P (normal temperature and pressure).

If the temperature is kept constant, the volume of a gas varies inversely with the pressure (Boyle's Law). Therefore, to express a volume of gas, measured at some other pressure, as its volume at 760 mm, the formula is

Volume
$$\times \frac{\text{Pressure (corrected)}}{760} = \text{Volume at 760 mm}$$

If the pressure on a gas remains the same and the temperature changes, the volume of the gas will vary directly ½712 (or 0 00367) of its volume at 0° C for each degree of change in temperature (Charles' or Gay Lussac's Law) The coefficient of expansion of nearly all gases is 0 00367

To convert the volume of a gas at temperature to C, and at constant pressure, to its volume at 0° C, the formula is

Volume
$$\times \frac{273}{273 + t^{\circ} C}$$
 = Volume at 0° C

The two conversion equations given may be combined into a single equation as follows

$$\frac{P \times V}{273 + t} = \frac{760 \times V_0}{273}$$

where P and V represent the experimentally determined pressure and volume, corrected if necessary, at the temperature t^p C, and V_0 represents the volume under standard conditions

If a volume of gas is in contact with water, evaporation continues until the tendency of the water to evaporate is equal to the tendency of its vapor to condence at which time the gas is said to be saturated with water vapor. The total pressure of the saturated gas and water vapor is

The realing on a brase-scale harometer is correct (x only one temperature an it rule corrected by the use of ac receivon table withen two table ight at which the cluin of mercury would stand if the temperature were 0° () for such a table refer to a handlock of chromatop or players. We also p 60%

equal to the sum of the pressures of the gas and of the water vapor (Dollon's Law). These are spoken of as partial pressures. The partial pressure of the water vapor increases with temperature. To determine the pressure of the gas in a dry state, the partial pressure of water vapor at the given temperature (see p. 709) must be deducted from the barometric pressure.

Example. The expired air of a man has a volume of 100 liters, the temperature in the gasometer is 22° C., and the brass scale barometer reads 767 mm. The pressure of expired air, dry, is:

The volume of dry air at 0° C. and 760 mm. is:

$$100 \times \frac{744.65}{760} \times \frac{273}{273 + 22} = 90.66$$
 liters

Equal volumes of different gases or vapors under like conditions of temperature and pressure contain the same number of molecules (Avogadro's Law). Thus the density or mass per volume of any gas or vapor will depend upon the weight of its molecule. The molecular weight in grams of any gas or vapor has a volume of practically 22.4 liters at 0° C. and 760 mm. Hence the density of 1 liter of a gas at 0° C. and 760 mm. is molecular weight + 22.4.

Density of Gases at 0° C. and 760 mm.

- 1 liter of oxygen weighs 1.4292 g.
- 1 g. of oxygen occupies $\frac{1}{1.1202} = 0.6997$ liter
- 1 liter of carbon dioxide weighs 1.9652 g.
- 1 g. of carbon dioxide occupies $\frac{1}{1.0659} = 0.5089$ liter

HEAT. The heat which we recognize by temperature is the energy of molecular motion. This property is imparted to matter by chemical action, electric currents, and mechanical work. In animals the chemical source only need be considered and we may confine our attention to that form of combustion in which the substance finally appears in the completely oxidized form.

In animal calorimetry the unit of heat is the large Calorie, abbreviated Cal., which is defined as the amount of heat necessary to raise the temperature of 1 liter of water from 15° C. to 10° C.

Heat of Combustion in a Calarimeter.

- 1 g. of hydrogen gas produces 34.5 Cal.
- 1 g. of charcoal produces 8.0 Cal.
- 1 g. of starch produces about 4.2 Cal.
- 1 g. of glucose produces about 3.74 Cal. 1 g. of sucrose produces about 3.96 Cal.

Heat of Combustion in Animals

1 g of average carbohydrate produces about 4 Cal

1 g of average fat produces about 9 Cal

1 g of average protein produces about 4 Cal

Oxidation of Carbohydrates. Carbohydrates, by the processes of digestion, undergo hydrolytic cleavage if necessary and are absorbed through the intestinal wall into the blood mainly in the form of glucose Under normal conditions the blood contains 0.1 per cent or less of this sugar in the free state, over 300 g or more may be stored as glycogen in the blood issues, principally in the muscles and liver Glycogen is readily reconverted to glucose and it serves as a deposit to be drawn upon in an emergency Glucose is the favorite body fuel, and is used produgally when the supply 19 plintiful and thriftily in periods of starvation

The oxidation of glucose is represented as follows

Carbohydrates contain hydrogen and oxygen in the same proportion as is found in water. When these substances burn, outside oxygen is used to unite with the carbon, forming a volume of carbon dioxide equal to the volume of oxygen absorbed. The ratio of the volume of earbon dioxide producted to the volume of oxygen absorbed is known as the respiratory quotient (R.Q.). This has a different value for each of the major food components and serves to determine what substances are being burned. From the above equation it is seen that the respiratory quotient for glucose is

 $\frac{CO_3}{O_3} = 10$ and that 1 liter of oxygen represents a liberation of 5 011 Cal In animal calorimetry the heat equivalent of 1 liter of oxygen is generally

accepted as 5017 Cal when earbohydrates are burned in the body Oxidation of Fats. The fats and oils of our food are largely mixtures of palimitin, stearin, and olem (see Chapter 3). These substances are of similar chemical composition and when they undergo oxidation they yield about the same amount of heat. After absorption, fat pusses from the blood to the treates where it is either burned or stored for future use. Fats have a high fuel value and they are continually being used by the body for this purpose. In the absence of carbohydrates, fats supply over 80 per cent of the body heat.

Fat combustion is usually represented by the oxidation of palmitin as follows

from the also exequation it is seen that when palmitin is harned the respirators question is 6.764 and that I liter of absorbed oxygen liberates

ANALYSIS OF THE OXIDATION OF MIXTURES OF CARBOHYDRATE AND FAT*
(Table of Zuntz and Schumburg modified by Lusk)

R O	Percentag Oxygen C	e of Total onsumed		e of Total roduced	Cai	per liter	of O•
n Q	Carbo- hydrate	Fat	Carbo- hydrate	Fat	Number	Log	Log + log 60
0 707	0	100 00	0	100 00	4 686	67080	44895
71	1 02	99 00	1 10	98 90	4 690	67114	44929
72	4 44	95 60	4 76	95 20	4 702	67228	45043
73	7 85	92 20	8 40	91 60	4 714	67342	45157
74	11 30	88 70	12 00	88 00	4 727	67456	45271
75	14 70	85 30	15 60	81 40	4 739	67569	45384
76	18 10	81 90	19 20	80 80	4 751	67682	45497
77	21 50	78 50	22 80	77 20	4 704	67794	45609
78	24 90	75 10	2ь 30	73 70	4 776	67906	45721
79	28 30	71 70	29 90	70 10	4 788	68018	45833
80	31 70	68 30	33 40	66 60	4 801	68129	45944
81	35 20	64 80	36 90	63 10	4 813	68241	46056
82	33 60	01 40	40 30	59 70	4 82a	68352	40167
83	42 00	58 00	43 80	ə6 20	4 838	08403	46278
84	45 40	54 60	47 20	52 80	4 850	68573	46388
85	48 80	51 20	50 70	49 30	4 862	68683	46498
80	52 20	47 80	54 10	45 90	4 875	68793	40603
87	55 60	44 40	57 50	42 50	4 887	68903	46718
88	59 00	41 00	60 80	39 20	4 899	69012	46827
89	62 50	37 50	64 20	35 80	4 911	69121	46936
90	65 90	34 10	67 50	32 50	4 924	69230	47045
91	69 30	30 70	70 80	29 20	4 936	69339	47154
92	72 70	27 30	74 10	25 90	4 948	69447	47262
93	76 10	23 90	77 40	22 60	4 961	69555	47370
94	79 50	20 50	80 70	19 30	4 973	69663	47478
95	82 00	17 10	84 00	16 00	4 985	69770	4758a
96	86 30	13 70	87 20	12 80	4 993	69877	47692
97	89 80	10 20	90 40	9 58	5 010	69984	47799
98	93 20	6 83	93 60	6 37	5 022	70091	47906
99	36 60	3 41	96 80	3 18	5 035	70197	48012
1 00	100 00	0	100 00	0	5 047	70303	48118

^{*}The last column has been added to facilitate the expression of oxygen absorbed per minute in terms of Calories per hour Characteristics are omitted (see last paragraph p 741)

4 655 Cal Zuntz and Schumburg old med slightly different figures for the oxidation of fat in the body. In their work the respiratory quotient was found to be 0 707 and the heat value for 1 liter of oxygen 1 656 Cal If only mixtures of carbohydrate and fat were oxidized, the respiratory quotients would vary between 0 707 and 1 00, and from their value it should be possible to determine the proportions of eight from their value it should be possible to determine the proportions of eight fittes, foodstuffs burned. A table analyzing the oxidation of such mixtures on the basis of the so-called nonprotein respiratory quotient was prepared by Zuntz and Schumburg and modified by Lust, (see p. 727).

Protein Metabolism. Proteins in the dict are completely broken down in the gastrointestinal tract to the form of amino acids, in which form they are absorbed into the blood and pass to the various parts of the body. Here they may be utilized either for incorporation into new tissue protein which is continually being broken down and resynthesized, or they may undergo metabolic reactions leading to the formation of nonprotein introgenous compounds of importance to the body, such as ereatine, thyroxine, adrenaline, etc Presumably they may also be utilized for the direct production of energy, although under ordinary circuinstances protein is not considered to be primarily a fuel Regardless of the intermediate steps in protein metabolism, which are discussed in detail in Chapter 33, the end products of protein breakdown in the animal body include not only carbon dioxide and water, as for carbohydrates and fats, but also the urea of the urine and certain other nitrogenous constituents of the unne and feces, as well as such compounds as unnary sulfate produced by the exidation of sulfur-containing amine acids. Hence to evaluate the contribution of protein to the total metabolism, and to distinguish it from the nonprotein metabolism the analysis of urine and feces is necessary in addition to the measurement of the respiratory gas exchance

A computation of protein metabolism by Locky is as follows 100 g of meat protein contains

 $52.38~{\rm g~C}$ $-7.27~{\rm g~H}$ $-22.68~{\rm g~O}$ $-16.65~{\rm g~N}$ $-1.02~{\rm g~S}$ of which there are climinated in the urine

9 406 g C 2 663 g H 14 099 g O 16 28 g \ 1 02 g S

769 g O

in the feces

1471 g C 0212 g H 0889 g O 037 g \

leavn g a residuum for the respiratory process of

41 o g C 44 ;

0961 L. II 769 g O

leaving

41 5 g C 3 439 g H for exclation

When CO2 is formed, 12 g C unite with 32 g O

Therefore 41 5 g C unite with
$$\frac{32 \times 415}{12}$$
 = 110 67 g O

When H2O is formed, 2 g H unite with 16 g O

Therefore 3 409 g H unite with
$$\frac{16 \times 3439}{2} = 27512$$
 g O

Total oxygen absorbed .

Total carbon dioxide produced

In the computation above, I g of urmary mitrogen represents

6 15 g protein × 4 25 (Cal) = 26 14 Cal

8 49 g O₂ × 0 6997 (hters) = 5 94 hters of O₂ for protein 9 35 g CO₂ × 0 5089 (hters) = 4 76 hters of CO₂ from protein

Respiratory quotient for meat protein
$$=\frac{476}{594}=0.801$$

Based on the analytical figures for the average protein it is estimated that 1 g of urnary nitrogen represents the metabolism of 6 25 g of protein, the absorption of 5 91 liters of ovygen, the production of 4 76 liters of carbon diovide, and the liberation of 26 51 Cal

An example of the practical application of the above constants is as follows

A man (aged 38.5 yrs, height 159 cm weight 90 lb) was tested 14 hours after the last meal. An analysis of the urine showed an exerction of 0.16 g of nitrogen per hour. The total respiratory exchange showed an absorption of 12.52 liters of oxygen and the production of 9.3 liters of carbon dioxide per hour.

6 16 g of urmary mitrogen represents

```
0.16 \times 6.25 = 1 g of protein metabolized with the absorption of 0.16 \times 5.91 = 0.95 liter of O_2, the production of 0.16 \times 4.76 = 0.76 liter of CO_2, and the liberation of 0.16 \times 20.51 = 4.24 Cal Total CO_1 (or O_2) = protein CO_2 (or O_2) = nonprotein CO_2 (or O_3) = 9.3 liters CO_1 = 0.75 liters O_2 = 8.54 liters CO_3 = 12.52 liters O_3 = 0.95 liter O_3 = 11.57 liters O_3
```

$$\frac{8.54 \text{ liters CO}_2}{11.57 \text{ liters O}_2} = 0.74 \text{ (nonprotein R Q)}$$

From Zuntz and Schumburg's table (modified b) Lusl, see p. 727) it is seen that when the nonprotein R Q is 0.74. I liter of O_1 represents the liberation of 4.727 Cal, and 12 per cent of the nonprotein heat comes from earholy drate and 88 per cent from fat

```
11 57 liters O<sub>2</sub> × 4 727 = 54 69 Cal (nonprotein)

12 per cent of 54 6) = 656 Cal from carbohydrate

88 per cent of 54 69 = 48 13 Cal from fat

54 69 + 4 24 (protein Cal) = total of 58 93 Cal per hour
```

In this computation 7 per cent of the body heat was derived from the combustion of protein, 11 per cent from carbohydrate, and 82 per cent from fat

As Richardson points out, the measured respiratory quotient is not necessarily the metabolic quotient and gives no information concerning the intermediate steps in the conversion of consumed O₂ to CO₂. Non-conductive processes which promote CO₂ production, such as acid formation (e.g., diabetes or excessive muscular activity) or aid retention (e.g., nephrits), and conditions which diminish the CO₂ output (e.g., insulin or alkali therapy, etc.) excresse an important influence on the respiratory quotient. Taking these factors into account, however, studies of conditions of abnormal carbohydrate and fat inetabolism such as diabetes and ketosis, and of the physiology of food utilization and muscular activity, have been greatly advanced by measurements of the respiratory quotient.

Respiration. The ventilation of the lung and the exchange of gases between the alveolar air and the blood are usually referred to as external respiration, as contrasted with the exchange of gases in the tissues, known as internal respiration (see Chapter 12). Any disproportion between external and internal respiration affects the oxygen or carbon dioxide content of the blood. The tension of oxygen in the blood is lowered when the lungs are extensively disordered, as in pneumonia, and during periods of supreme muscular effort, under these conditions the oxygen requirement of the tissues is greater than the oxygenating power of the lung An unusual increase in the supply of oxygen in the lung has but a very slight effect on the tension of oxygen in the blood, and practically no effect on the rate of combustion in the tissues. Of greater interest is the carbon dioxide tension of the blood, which is nicely regulated by an adjustment of the ventilation of the lung, and of the eirculation of the blood, so that these mechanisms parallel the rate of combustion in the tissues.

There is a general parallelism between the heat production, the heart rate, and the ventilation rate However, the heart rate is not solely determined by the requirements of respiration, and ventilation can be voluntarily controlled A rapid heart and an increased ventilation rate make one suspect that the metabolism is increased, but the rate of heat pro-

duction cannot be predicted from these phenomena

EXTERNAL RESPIRATION Atmospheric air is of very constant composition the world over Advantage is taken of this fact in computing the respiratory exchange, that is, the amount of oxygen absorbed and the amount of carbon doxide produced in a given time

Atmospheric air has the following composition

Carbon dioxide Oxygen Nitrogen 0 03 per cent

20 94 per cent 79 03 per cent

Under ordinary conditions, at each breath a person inspires about 500 ml of air and of this amount only 330 ml actually enter the lung and diffuse more or less with the 3000 ml already there. The expired air includes the 140 ml of the "dead space" and 350 ml of partially mixed air.

² For further discussion of the gas exchange in blood and tissues see Chapter 24.

from the lung (see Fig. 195). Expired air contains about 16 per cent oxygen, 4 per cent carbon dioxide, and 80 per cent nitrogen

If we exhale forcibly and collect the last portion, the so-called alveolar air, we find that it contains about 14 5 per cent of oxygen and 5 6 per cent

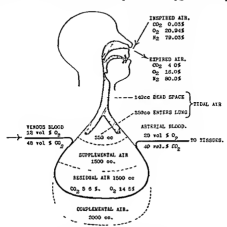


FIG 195 INSPIRED AND EXPIRED AIR

The tidal air is that which enters and leaves the body with each normal respiration. The supplemental air is that which can be foreibly expired after a normal tidal expiration, this leaves in the lungs the residual air which cannot be voluntarily expelled. The complemental air is that which can be foreibly drawn into the lungs over and above a normal tidal inspiration. The vital capacity is the greatest volume of air which can be expired following a forced inspiration, it includes the tidal, the supplemental and the complemental volumes.

The inspired or atmospheric air is of constant composition. The composition of the expired air is not constant, it is determined by the varying factors of combustion in the tissues the rate of blood flow, and the rate of ventiation in the lung. The already air is the last portion of the supplemental air to be expired and represents the composition of the residual air, and its gases are in equilibrium with the arternal blood. The blood in its passage through the lung gives off carbon dioxide and absorbs oxygen.

of carbon dioxide. That is, the air deep in the lung, where it comes in close proximity to the blood, has the greatest concentration of carbon dioxide and the lowest concentration of oxygen. These gases are in equilibrium with the gases in the arterial blood.

Fig. 195 shows that after an ordinary expiration, 1500 ml. of additional air can be fortably expired; then, if the greatest possible inhalation is made, almost 4 liters of fresh air can be drawn into the lungs. This greatly dilutes the carbon dioxide in the alweolar air and induces a rapid passage of the gas from the blood. If forced breathing is continued, a large amount of carbon dioxide, which is ordinarily stored in the blood, is in this manner "washed out" and may be erroneously regarded as carbon dioxide produced during a period of a metabolism test.

The increased supply of oxygen due to forced ventilation has no effect on metabolism tests; in fact, as bas previously been stated, one can breathe pure oxygen without appreciably increasing its absorption. This fact is taken advantage of in the "closed circuit" types of respiration anparatus which require the rebreathing of air rich in oxygen

Nitrogen plays a passive role in respiration; the amount expired is the same as the amount inspired. As there is a definite proportion between the oxygen and the nitrogen of the atmosphere, it is possible to calculate the amount of oxygen inspired on the basis of the nitrogen in the expired air. The formula is:

$$V = \frac{20.94 \times \text{per cent N}_2 \text{ in expired air}}{79.03}$$

when V = volume of oxygen in inspired air corresponding to 100 volumes of expired air.

It must be emphasized that in order to measure successfully the carbon dioxide produced during a metabolism test of short duration, rentilation must be on involuntary act

Energy Requirement. The energy requirement of the animal body may be divided into two functional classifications, viz, that for basal metabolism and that for active work Basal metabolism includes the energy expended in respiration, blood circulation, intestinal contractions, activities of various organs, maintenance of muscular tonus, thermal equilibrium, etc. The basal metabolic rate is influenced by the amount of active protoplasmic mass (bence by height, weight, surface area, agsex, composition of the tissues, etc.) and is governed by endocrine organish particularly the thyroid and pituitary glands. For a discussion of the clinical aspects of basal metabolism and its measurement, see p. 734.

The energy consumed in work, play, and indeed all forms of voluntary activity, imposes an additional requirement for fuel over the basal, which depends on the nature and extent of the muscular work. Whereas an average man expends about 100 Cal per hour while sitting at rest, his metabolism may increase to as high as six times this value with extreme physical effort.

Mary Swartz Rose summarized the hourly expenditure of Calories of an average 70-kg man under various conditions as follows: sleeping 65, awake lying still 77, sitting at rest 100, standing relaxed 105, valking slowly to moderately fast 200-300, running 53 miles per hour 570, swimming 500, walking down stairs 364, walking up stairs 1100, "light exercise" 170, "active exercise" 290, "very severe exercise" 600, dishwashing the stairs 1100, "light exercise" 170, "active exercise" 290, "very severe exercise" 600, dishwashing the stairs 1100, "light exercise" 170, "active exercise" 290, "very severe exercise" 600, dishwashing the stairs 1100, "light exercise" 1100, stairs 1100,

144, earpentry or painting 240, sawing wood 480. From estimates such as these one may predict the calorific content of the diet necessary to meet individual requirements. An example for a 70-kg. man is given by Sherman as follows:

8 hrs sleep at 65 Cal. = 520 Cal. 2 hrs. light exercise at 170 Cal. = 340 Cal 8 hrs carpenter work at 240 Cal. = 1920 Cal. 600 Cal. Total food requirement for the day, 3380 Cal.

The total energy requirement of different types of workers ranges from a minimum of 2000 to 2500 Cal. per day (white-collar workers) to a maximum of 4000 to 6000 (lumbermen, exeavators, etc.), of which about 1400 to 1900 Cal. are consumed in basal metabolism and the balance in various forms of activity.

Food calories consumed in eveess of an individual's energy requirements are deposited principally in the form of adipose tissue which is drawn on for energy when the dietary intake of calories falls below the expended level. A deficit of approximately 3500 calories is required to deplete the body of one pound of adipose tissue When the fat depots are exhausted, as in acute inanition, muscle protein is used as a source of energy.

BASAL METABOLISM

Basal metabolism, or the basal metabolic rate, is an expression of the heat production of the body in complete mental and physical repose, and in the postabsorptive state. For a successful test the subject must be in a "eomfortable" environment, if "too hot" or "too cold" the result will be abnormally high. The body temperature must be within the normal limits. DuBois estimates an increase in metabolism of 7.2 per cent for each degree Fahrenheit in fevers. The patient must be quiet and preferably experienced with the test. Mental activity alone has but a slightly elevating effect upon the heat production. Emotional excitation will raise the metabolism 20 per cent or more probably owing to the stimulation of the adrenal glands. The emotional factor constitutes the greatest source of error in hasal metabolism determinations. It is not difficult to insure muscular repose and physical comfort, but it requires the utmost sympathy and tact to allay the apprehensions of the patient and insure physiological repose. A common practice is to test the patient repeatedly and to accept the lowest or the last result as the true basal rate; the fallacy of this is obvious when one considers that the number of such tests is usually determined by the endurance of the patient. DuBois conducts three or four tests and takes the average of the two lowest that show fair agreement. Even under favorable conditions it is possible, in unstable individuals, to get variations as high as 6 per cent in as many hours. These variations are physiological, not technical; the technique is more accurate than our ability to recognize or control emotional changes.

Going to and from work, for example.

Clinical Interpretation of Basal Metabolism. In 1893 the clinical use of calorimetry was presaged by Friedrich Muller, who observed that patients with Graves's disease lost weight and had a marked introgenous waste despite the fact that the diet was adequate to maintain a normal state of nutrition. This observation was soon verified by Viagnus-Levy, who determined the now well known action of the thyroid gland in regulating the rate of combustion in the body. It is true that other factors affect the rate of heat production and that certain well recognized diseases are accompanied by metabolic changes, but in the mojority of cases ioria tions in the basal metabolism can be interpreted as ioriotions in the function of the thyroid gland.

In an extensive study of basal metabolic rate determinations, Boothby and Sandiford showed that 921 per cent of normal individuals have a basal metabolic rate within ±10 per cent, and 93 per cent within ±10 per cent of the DuBois standards. They found that a smaller percentage of these same subjects bad hasal metabolic rates within the same limits when the Harris and Benefite standards were used. These standards are

discussed in the following section

In clinical cases the basal metabolism may vary from 40 per cent below to 130 per cent above the average normal. In this discussion a tabulation of the findings in various disorders is avoided because standing alone the values are misleading. If uo other cause can be found for an abnormal heat production the result may be cautiously attributed to a disordered activity of the thyroid gland. Glands other than the thyroid may affect the rate of cellular combustion but at the present time we know of only two body substances which change the rate of heat production namely thyroine and adrenaline. The latter has a rapid but fleeting stimulating effect, the effect of the former lasts for several weeks. When the activity of the adrenal glands is decreased as in Addison's disease the basal metabolism is found to be subnormal. Hyperactivity of these glands may account for the temporary increase in metabolism cocasionally found in nervous patients who also show corresponding variations in pulse rate and in blood pressure.

The basal metabolic rate does not establish a diagnosis of byperthyroid ism \(^1\) patient afflicted with byperthyroidsm may be tested during an intermission and the basal metabolism would be within the normal limits despite the persistence of such chinical signs as nervousness palpitation tremors and exophthalimos Converely the test may detect active hyperthyroidism in patients showing few of the classical symptoms of the disease. The test does not tell the surgeon when it is safe to operate upon the patient it informs him when it is least dangerous. He must hase his operative prognosis upon the clinical condition of the patient and not upon the basal metabolic rate.

The order of hyperthyroidism is accompanied by a progressive increase in heat production which may reach over 100 per cent above the average normal and per ist for weeks after the glandular activity has subsided Therefore repetited tests are necessary in order to follow the course of the disease an increasing rate indicates an active state of the gland and

naturally precludes operative intervention, a decreasing rate gives a more hopeful outlook

The pulse rate is largely dependent upon the rate of metabolism Chinically, a decreasing pulse rate indicates a subsidence of the hyperthyrodism. Few patients with pulse rates below 50 have a basal metabolic rate above the average normal and few patients with pulse rates below 85 have any considerable increase in their metabolism.

Heat comes from oudation in the active protoplasmic tissue of the hody. The rate of heat production is not determined by surface cooling, as one method of measurement would seem to imply (see footnote 17, p. 738), in reality, heat dissipation is regulated according to heat production. Any rapid change in the relative amount of protoplasmic tissue will cause changes in the basal metaholic rate when expressed in terms of weight or surface area. This partity explains the gradual drop in the basal metaholic rate during the glandular and muscular waste of starvation and in bospital, patients during their early confinement to bed. A rapid accumulation of fluid in the body has a similar effect and it is quite common to find nephritics with marked edema having a basal metabolic rate 20 to 30 per cent below the average normal. This must not be interpreted as a sign of hypothy roidism.

Under the hasal conditions we measure the heat resulting from activities of the vital organs and from the intracellular chemical changes associated with life, growth, and development. These facts must be borne in mind and due allowance made for any unusual activity before the results are attributed entirely to the thyroid gland. Dyspnea, bypertension, cardiac decompensation, and tremors involve increased muscular activity and so raise the heat production. Certain diseases are characterized by increased cellular activity and in these cases the heat production is increased The most outstanding of this group is leukemia, in which disease the basal metaholic rate may he as high as in severe Graves's disease Polycythemia, anemia, and Paget's disease of the bones are other conditions in which the increased heat production probably results from increased cellular activity. In this group possibly belongs the acute stage of acromegaly. In these conditions the hasal metaholism is rarely over 25 per cent above the average normal. The possibility of selfadministration of drugs such as iodine or thyroid extract must be considered in the interpretation of hasal metabolic rate

From hirth to the age of one and a half years the basal metabolism increases at a remarkable rate. This is followed by a gradual decline until full growth and development are attained, constancy characterizes the rate in adult life, with a slight decline as old age advances. Variations in the rate of growth and development in childhood cause abnormal results

when judged by the age standard In prematurely developed children the basal metabolic rate is low according to the age standard, but is probably normal for the stage of development if this could be accurately expressed The difficulty in applying the test to children has resulted in a paucity of normal tests and great variation in the results obtained The normal biological variations are much greater in children than in adults and may be greater than the pathological changes anticipated by the physician Fortunately the test is rarely required before the age of puberty and the normal standards beyond that age are fairly well established

The metabolism of women averages about 12 per cent below that of men, and owing to the menstrual cycle, is more variable. There is usually, though not invariably, a premenstrual rise and a postmenstrual fall in the basal metabolic rate, which should be considered in the interpretation

There is some evidence of lower metabolic rates among certain oriental and tropical races

STANDARDS OF NORMAL BASAL METABOLISM AND CALCULATIONS & The heat produc tion of normal individuals under hasal conditions largely depends upon the factors of age height and weight The normal standards are based upon thousands of deter minations in several centers of investigation. A better appreciation of basal conditions and the climination of a high proportion of first tests has resulted in a tendency toward lower standards in recent years. At the present time there are three different systems of predicting the normal heat production

1 jub and DuBous determine the heat production in relation to the surface area of the body, the surface area is estimated by the DuBois and DuBois' formula which is based on the height and weight

$$A = We^{in} \times He^{in} \times 71.84$$

where A equals the area in sq em Wt the weight in kg and Ht the height in cm In routine work the nomogram of Boothby and Sandiford* (Fig. 196) may be used for determining the surface area Stoners has computed tables of values for this formula in unit cm and kg intervals for the height range of 110 to 200 cm and the weight range of 20 to 110 kg.

The original so-called Sage standards of Aub and DuBois have been modified on the lass of larger numbers of normal controls Bailey sis table covers the ages from 4 to 65 in yearly intervals while Boothby and Sandifordii give values by years from 5 to 19 and in 5-year intervals between 20 and 79 (see also table on p 738) These standards fail to reflect the modern downward trend but since they are based on lests on patients they may be better suited for clinical purposes than lower standards l ased on trained subjects

2 The Harris-Benedict¹³ multiple prediction equations and tables¹³ are based on

For prediction tables of normal heat production reference should be made to books deal ng in this subject (see the Bibliograph) at the end of this chapter)

Aub and Dulion Arch Internal Med 19 823 (1917)

^{*}Ault and Dubble a real reserves sees 1's acc (1917)

*Boothy and Sandford Besters Med Sur 7 565 (1921)

*Stoner J Lab C 1'n Med 11 325 (1925)

*Stoner J Lab C 1'n Med 11 325 (1926)

*Bladey J Lab C 1'n Med 4 657 (1921)

*Bloothly and Sandford daw J Playmed 200 (1979)

*See also Krogh's table 1'n

*Polyton Bester Webelium to Hessik and Disease 3'd ed 11 lidelejbin Lea & Feb ger 1936.

Harris and Benedict Carnegie Inst Bash Iub to 279 1919 *Tables with greatly simplify the use of these formulas may be found in Carpenter Caracque Inst. Wash. Pub. Vo. 303, 1921. See also DuBois a book cited above.

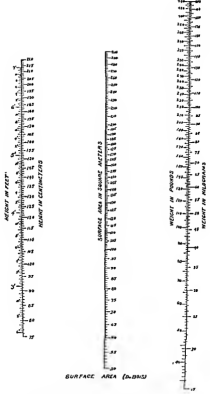


Fig. 196. Surface-Area Nonogram,
A line connecting the height with the weight intersects the
middle line at the corresponding surface area
Copyright 1920 by Boothby and Sandford.

a statistical study of the available data for the basal metabolism of normal men and women. The equations are, $\,$

For men,
$$H = 664730 + 137516H + 50033S - 67550A$$

For women, $H = 6550935 + 95634H + 18496S - 46756A$

in which H = total heat production per 24 hours, W = weight in kilograms, S = stature in centimeters, and A = age in years. The Harris-Benedict standards a verage about 3 to 4 per cent lower than the Sage standards, and according to Benedict¹⁴ they should be even lower, especially for women. He recommends lowering the standards for women by 5 per cent. Benedict calls attention to the distinction between "hospital normals" and physiological standards, the use of the former being justified for chinical purposes only.

STANDARD VALUES FOR MEAN LINERGY PRODUCTION IN RELATION TO AGE AND SEX*

lge at Last Birthday	Calones ner	sį m per hr
Year	Males	Females.
6	53 0	50 6
7	52 5	49 1
8	51 8	47 0
9	50 5	45 9
10	48 5	45.8
11	47 2	45 3
12	46 8	44 3
14	46 4	41.5
16	45.7	38 9
18	43 3	37 0
20	41 8	36 2
25	40 4	35 9
30	39 6	35 8
35	38 9	35.7
40	38 2	35 Q
50	37 0	34 5
60	35 8	33 0

The method of predicting the normal heat production, and the normal standards, for infants and children are the subjects of several papers 12

THE RESPIRATORY QUOTIENT (R.Q.) This is the ratio of the volume of carbon dioxide produced to the volume of oxygen consumed during the same time interval

R Q =
$$\frac{\text{CO}_2 \text{ produced}}{\text{O}_2 \text{ consumed}}$$
 = $\frac{\text{volume CO}_2 \text{ in expired air} - \text{volume CO}_2 \text{ in expired air}}{\text{volume O}_2 \text{ in inspired air}}$ = $\frac{\text{volume CO}_2 \text{ in expired air}}{\text{volume O}_2 \text{ in expired air}}$

The measurement of the volume of inspired air is attended with technical difficulties. However, the actual volumes of gas disappear from the working formula as demonstrated below. If we let

 V_r and V_s = volume of expired and inspired air C_r and C_s = per cent CO_2 in expired and inspired air O_r and O_s = per cent O_s in expired and inspired air V_s and V_s = per cent V_s in expired and inspired air.

we may rewrite the above equation as follows

$$R Q = \frac{C_{\bullet}V_{\bullet} - C_{\bullet}V_{\bullet}}{Q V_{\bullet} - Q V_{\bullet}}$$
 (1)

Since nitrogen is neither absorbed nor evolved its volume in inspired and expired air remains unchanged. Therefore its percentage in inspired and expired air is in inverse ratio to their respective joilmes. That is

$$N, N, = V, V$$

or

$$V_{\bullet} = \frac{N_{\bullet}V_{\bullet}}{N_{\bullet}}$$

Substituting this value in (1) and simplifying we get

$$RQ = \frac{C_t \setminus_1 - C_t \setminus_2}{C_t \setminus_1}$$
 (2)

Substituting the values given for the composition of atmospheric air (p. 730) for O_{ij} , C_{ij} and A_{ij} , we obtain

$$RQ_{*} = \frac{7903C_{*} - 003V_{*}}{2003W_{*} - 7903O}$$
 (3)

Simplifying

$$RQ = \frac{C_e - 0.00038 V_e}{0.265 V_e - Q_e}$$

Since 0 00038N, never varies significantly from 0 03, we have finally

$$R Q = \frac{C_{\bullet} - 0.03}{0.265 \, V_{\bullet} - O_{\bullet}} \tag{4}$$

From this equation it is possible to calculate R Q, simply from the CO₂ and O₂ percentages in expired air, since A₂ is obtained by difference

BASAL METABOLISM OR BASAL METABOLIC RATE (B M R.) This may be defined as the percentile variation of the observed from the normal or predicted heat production for an individual of given highly weight, age, and sex in a postabsorptive state and in complete mental and physical repose. The best production for a given period is

¹⁸ Talbot Physiol Rets 5 477 (1925) Stark 4m. J Physiol , 111, 630 (1935), Talbot, Wilson and Worcester 1m. J Discoses Child , 53, 273 (1937) Lewis kinsman, and Iliff Am. J Discoses Child , 53, 345 (1937).

obtained by multiplying the volume, at normal temperature and pressure, of oxygen consumed during that period by the calorific value for oxygen corresponding to the observed (or assumed) R Q Basal heat production may be expressed on an honrly or daily basis. The general formula for I asal metabolic rate is therefore

In this equation, p is the barometric pressure in mm of mercury corrected for tension of aqueous vapor, s the observed volume, in liters, of oxygen consumed during the basal test period, m the duration of the hasal test period in minutes, t the absolute temperature (° C + 273), C the calorific value per liter of oxygen corresponding to the observed (or assumed) respiratory quotient (see the table on p 727, in the oxygen consumption methods the R Q, is assumed to be 0 82, which corresponds to a calorific value per liter of oxygen of 4 825), and N, the normal or predicted basaf heat production obtained from one of the various tables of standards

In the usual form of oxygen consumption apparatus equipped with volumetre scales and CO, absorbents, v in equation (5) is obtained by subtracting the reading at the end from that at the beginning of the test period

When the gasometric method is employed,

v = volume O, in inspired air - volume O, in expired air or

$$\sigma = 0.010 \epsilon V_4 - 0.010 \epsilon V_4 \tag{6}$$

By substituting the value for V_i as above, and simplifying

$$v = 0.01V_{\bullet} \left(O_{i} \frac{N_{\bullet}}{V_{i}} - O_{\bullet} \right)$$

in which V, is the gasometer reading R multiplied by the gasometer factor, f, hence the volume of expired air in liters. By further substitution of the values for Oi and Ai given above, and simplifying

$$v = 0.01Rf(0.265N_{*} - O_{*})$$

in which the variables are R N. and O. This expression for v may be incorporated in equation (5)

The Aub and DuBois tables give the normal heat production in Cal per sq m of body surface per hr (A) which when multiplied by the surface area in sq m (S) gives the predicted hourly heat production. Making the proper substitutions in the above equation and collecting constants we obtain the following simplified formula "

B VI R. =
$$\frac{2155peV}{tmSA}$$
 - 100

In the Harris and Benedict tables for normal heat production the values are expressed on a 24-hour basis. The coefficient in the equation becomes therefore 24 × 2155 = 51720, and the equation is

exchange, as well as data cards which are very useful in routine practice

According to Magnus-Levy the metabolism of protein furnishes on the average 15 per cent of the total calories. This may be taken into account by using figures for the calorific while of lifter the form the same way be taken into account by using ngures for the carbon of lifter the form of the same same than those given in the table on p 727 as Stoner (Roferon Viel & Sury J 182, 198 (1923) J Leb Clin. Wed. 12, 884 (1927) 184 (1927)) has described been ad other simplifications in the calculations for respiratory

$$B M.R = \frac{51720 prC}{tmH} - 100$$

in which H represents the predicted 24-hour heat production

In the oxygen consumption methods R Q is assuined to be 0.82. Hence 4.825 is substituted for C, in the above equations (see the table on p. 727) and the coefficients become 10,400 and 250,000 respectively. Further simplification is attained in routine RESPRATION LABORATORY.

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TIG 197 FORM SHEET FOR STEDY OF RESPIRATORY METABOLISM

practice by standardizing the basal test period and merging the value for m into the coefficient

The calculations may be performed either on calculating machines or logarithmically, in which case fix-place tables are used and the characteristics omitted, since the decimal place may be pointed off in the final result by inspection. A form sheet for logarithmic computation described by Boothby and Sandiford is shown in Fig. 197

The calculations involved in the interpretation of Lymographic records of respiratory exchange are described in the legend to Fig 200

HUMAN AND ANIMAL CALORIMETRY The rate of combustion may be determined by directly measuring the heat given off by the body Livery animal has an optimum tem perature and this temperature is maintained by a nice balance hetween the heat produced by the life processes and the heat lost by the organism 21 Gephart and DuBois estimate that in man 24 per cent of the total heat loss results from the evaporation of water from the lungs and skin A small amount of heat is lost in warming the ingested food the urine the feces and at times the body itself, the remainder is lost by con duction and by radiation

Lavoisier and Laplace measured the heat loss by placing a guinea pig in an ice chamber for ten hours and observing the amount of ice melted. Over a period of many years, the elaboration of this simple form of calorimeter has culminated in the con struction of the Atwater-Rosa-Benedict Respiration Calorimeter 22 With this app3 ratus the heat of the body is determined indirectly from the respiratory exchange and directly by a careful computation based on observations of all the known means of heat loss

The principle of this apparatus was applied by Armsby22 in the claboration of 8 calorimeter for use with farm animals. Benedicta and associates also devised a respira tion chamber for use with domestic animals. This apparatus was later modified for use in human calorimetry 16 Respiration calorimeters have provided the means of acquiring most of the present-day knowledge of energy metabolism. They have been extremely valuable in proving the accuracy of indirect calorimetry

Because of the expense of installation and the technical difficulties of operation of respiration calorimeters heat production is usually measured indirectly 18 from the gaseous exchange Two types of methods are in use (1) Open-circuit methods in which atmospheric air is breathed and expired air collected and analyzed, and (2) closed curcuit methods in which oxygen-cariched air is breathed and the consumed oxygen measured The former procedure provides a more complete picture of respiratory exchange since it permits determination of the respiratory quotient On the other hand the closed excust methods are much simpler technically and where only total metabolism must be determined are equally satisfactory in the closed circuit method an average respiratory quotient of 0.82 corresponding to a calorific value for O1 of 4 825 Cal per later 18 assumed

The various types of respiration apparatus on the market are simplified modifications of Dr F G Benedict s76 clinical respiration chamber

Basal metabolism is usually determined in 10- to 15-minute periods using either one of the Benedict closed-circuit methods as developed in the Carnegie Nutrition Laboratory or a modified Tissot or open-circuit method. Obviously the determination of urmary nitrogen in such tests is impracticable. It has been the custom to apply the calorific values in the Zuntz and Schumburg table (see p. 727) directly to the respira tory exchange as measured without computing separately the protein metabolized DuBois points out that this procedure grace results about 1 per cent too high

The example described below serves to illustrate the hasic principles employed in

¹¹ For a review see DuBois Harry Lect 34, 88 (1939)

¹² Atwater and Ross Report of Surve Agric Exp. Sta 1897 p 212 Atwater and Benedict Caraçue Ind Wash Fub No 42 1905

Benedict Caraçue Ind Wash Fub No 42 1905

Armsby and Free Bull 51 U S Dept of Agriculture (Bureau of Animal Industry)

²⁴ Benedict Coropatchinsky and Ritzman Abderhalden a Handbuch der biologischen

Arbeitsmethoden 4 (part 13) 619 1934 15 Newburgh Johnston Waley Sheldon and Murrill J Nutration 13 193 (1937)

²⁶ Later developments in closed-circuit apparatus and comparison with open-circuit methods are discussed by Benedict Boston Ved Surg J, 193, 507 (1920)

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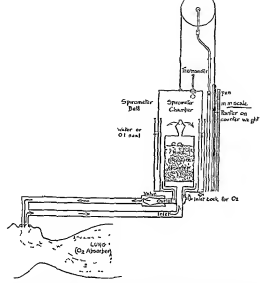


FIG 198 DIAGRAM OF THE BENEDICT-ROTH RESPIRATION APPARATUS

The patient breathes through a monthpiece The nose is clamped Two tubes connect the mouthpiece with the spirometer. An inspiratory Said valve directs the oxygen from the spirometer chamber to the patient an expiratory valve insures its continuous circulation. The expirations pass through soda lime which removes carbon dioxide. The calibrated spirometer bell is balanced by a counterweight which carries a pointer. The pointer shows the movements of the bell on a millimeter scale. A test is started with the chamber full of oxygen as oxygen is absorbed by the patient the volume in the chamber decreases. The loss of volume represents oxygen absorbed during the period of the test. Correction is made for any change in chamber temperature during the test.

Slightly modified from Roth Boston Med Surg J 186, 491 (1922)

the oxygen-consumption type of measurement. Specific directions are furnished with various commercial instruments adapted for office or hospital use

The Benedict-Roth respiration apparatus (Figs. 193, 199) is extensively used in chaics for determining the basal metabolic rate of patients. Roth¹⁷ has dispensed with the

²⁷ Roth Hoston Ved Surg J 188, 457 491 (1922) This and other types of basal metabolism apparatus for climeta use are nade by Warren E Collus Inc. 553H Huntington Ne Boston Mass. Sunborn Co. 39 Osborn St. Cambri Ige. Mass. and Jones Metabolic Equipment, Co. 1870 Ogden Ave. Cl. rago Ill.

electric blower of the Benedict machines and has inserted two Saidy valves to insure circulation of the oxygen. The spirometer bell is designed to have a volume of 20.73 ml for every millimeter of height. This greath simplifies the calculations. The fall of the spirometer bell during a test of 6 minutes represents the volume of oxygen al sorbed by the patient. As with other Leachdet appractant of a respiratory quotient is assured to be 0.82, this gives a bart value for 1 liter of oxygen of 4.82. Cal. "With a bell of this size, each min. of the fall of the bell in a six minute period represents exactly 1. Cal.

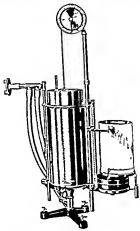


Fig 199 Benedict Roth Respiration Apparatus with Kymograph.

per hour (1 mm in 6 inim = 2073 inl = 0 2073 liter per hour 0 2073 × 4.825 = 1 Cal per hour)

The form of this apparatus illustrated in Fig. 199 is provided with a kymograph which is self tune-marking in minute periods. A pen attached to the counterweight traces the reputatory excursions and shows the fowering of the sprometer bell due to the alsorption of oxygon. Fig. 200 illustrates the method of drawing the "oxygen casumitton line and of measuring the rise of this line for any bermitted period which may be selected on the tracing. If the moniture content of the circulating air is high as is the case, when Wilson sods line is used as an absorber a correction must be made for water vapor. Roth²⁰ gives the factors for reducing the volume of

³⁶ A table f r reducing volumes of 80 per cent saturated air to 0° C and 760 mm dry is given by it it. Boston Med. Surg. J. 184, 4.7 (1922).

absorbed oxygen to 0° C, 760 mm, dry (when the vapor tension is 80 per cent of saturation)

Procedure. The patient is tested before breakfast and after a 12- to 15-hour fast A record is made of the sex, age, height, and weight. A rest in a semi-reclining position for half an hour must precede the test. During the rest period, the pulse rate, the respiration rate and the temperature are observed, and the patient is briefly instructed in order to secure his confidence.

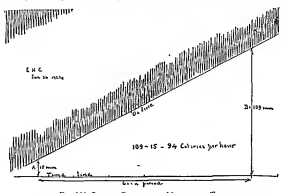


FIG 200 GRAPHIC RECORD OF A METABOLISM TEST

E H C, male, aged 38 Height 176 cm Weight, S4 kg Surface area, 20 sq m Barometric pressure, 745 mm Average temperature, 24° C (no merease during test) Rise of O₂ line in 6 mmutes 94 mm = 94 Cal per hour Correction for temperature, pressure, and water vapor 94 × 0 879 = 82 63 Cal per hour Divide by surface area \$2.63 - 2 = 413 Cal per square meter of surface per hour, which is 5 per cent above the average normal (39 5) for this individual (Roth)

For an example of a simplified data card for taking records of instruments not equipped with hymographs and for making the calculations, see Stoner Boston $Med\ Surg\ J$, 189, 195 (1923)

and cooperation. The apparatus is tested for leaks as follows: a stopper is placed in the opening of the mouthpiece, a 200-g weight is placed on the bell. The pointer should remain stationary for a minute or more. The weight and stopper are removed and the spirometer is filled with oxygen. The mouthpiece and later the nose clip are applied to the patient, care heing taken to avoid leaks. The kymograph²³ is started. The temperature of the spirometer, and the harometric pressure are recorded. The test is continued until a

[&]quot;In the case of sprometers not equipped with ky mographs, readings on the scale are made at the beginning and end of the periods Stoner (Boston Vict Surg J. 188, 193 (1923)) divides a 10-minute period into fix 2 minute periods taking the average of 10 sprometer readings at the beginning of each period in this way he is able to select a 4 minute period showing the least ony gen consumption, and also to avoid the error due to wide fluctuations in mid-vidual respirations.

satisfactory uninterrupted section of exactly 6 minutes can be selected for subsequent measurement. Before stopping the test, a weight of 50 g. is placed on the bell and operation is continued for a few minutes; if a leak occurs it is shown by a sharp rise in the oxygen consumption line. The temperature of the spirometer is again recorded. Fig. 200 shows a graphic record of a test, and the legend gives the necessary calculations for determining the basal metabolic rate.

Wesson 20 has described an apparatus and procedure for determining the respiratory quotient in small animals. A multiple-chamber respiration apparatus for small animals, based on the oxygen-consumption method, has been described by Benedict.21

For research purposes the expired air may be collected in a gasometer or in an impervious rubber or plastic bag and subjected to analytical determination of CO2 and O2 content. Methods hased on the use of the Haldanc-Henderson gas-analysis apparatus are employed. These procedures offer greater flexibility and more information but require more expensive equipment and a high degree of manipulative skill.

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^{**} Wessen: J Bud Chem , 71, 499 (1927)

is Henedict J. Valration 3, 161 (1930). This apparatus can be secured from Warren E. Collins, Inc., 55311 Huntington Ave., Boston, Mass.

Hormones

The endocrinc organs, or glands of internal secretion (see Fig. 201), are tissues whose function it is to secrete certain specific chemical substances known as hormones into the blood stream, which distributes them to all parts of the body. Certain tissues respond to their presence in a characteristic way, increasing, decreasing, or modifying their processes of growth or of metabolic or physiological activity. The hormones are therefore chemical messengers serving to integrate the various activities of the

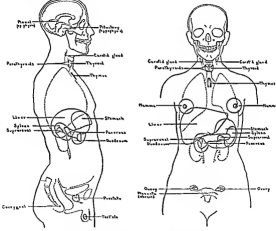


Fig 201. SCHEMATIC CHART OF ENDOCRINE STREEM.
From Barket, et al. Endocrinology and Metabolum, New York, Appleton, 1924

hody; they may themselves undergo metaholic transformations, such as partial or complete oxidation, or reduction; and recognizable products of these transformations, or the hormones themselves, may be exercted in the uring and in the bile.

The secretory activity of the endocrine organs varies from time to time As the chart on p. 750 shows, the morphological structure and the endocrine activity of the gonads, adrenal cortex, and thyroid are largely controlled by hormones produced by the anterior lobe of the pituitary

(anterior hypophysis) which also produces hormones acting directly on the structure and function of nonendocrine tissues. The anterior pituitary in turn is partly controlled (directly or indirectly) by the concentration of hormones produced by the gonads, thyroid, etc., and also by the nervous system directly Similarly, the posterior pituitary and the adread medulla are subject to direct nervous regulation. The control of the endocrine activity of the panereas, parathyroids, and other possible sources of hormones is not fully understood. It will be noted in the chart that a single organ, or a single metaholic process, may be affected by more than one hormone, and often also by factors not endocrine in nature.

The hormones do not vary significantly in chemical structure from species to species, hence extracts from the endoerine organs of animals may often he used in treating disorders due to insufficient activity of a human gland, synthetic or artificial products, allied to or identical with the natural hormonics, may be less costly. Disorders due to excessive activity—for example, when a tumor develops from endocrine cells—are often treated by surgical removal of the abnormal tussue, or, more rarely, by selective inhibitory drugs or hy irradiation with x rays. Chemically, the known hormones may be divided into two main groups. The first may be referred to as introgenous, and range in size and complexity from simple bases such as adrenaline to proteins of high molecular weight and undetermined structure. The second group, more homogeneous, may be called the storoid hormones.

THE STEROID HORMONES OF THE GONADS AND ADRENAL CORTEX

General Chemistry of the Steroids and Steroid Hormones The basic saturated carbon ring skeleton of the steroids is illustrated in Formula 1

Carbon Ring Skeleton of the Steroids (1)

Depending on the chemical nature of the substituent (R) at C_{17} , the steroids may be divided into five classes

(a) The sterols like cholesterol and ergosterol (see Chapters 11 and 35), where R consists of an eight- or nine-earbon atom alphanic side chain

30), where it consists of an eight-or nine-carbon atom alphatic side chain (b) The bile acids (p 110) where R consists of a five-carbon side chain terminating in a carboxylic acid group

(c) The cardiac aglycones breakdown products of the heart-stimulating glycoudes of Strophanthus Digitalis etc, which are characterized by a lattone ring substituted at Car.

(d) The sapogenins derived from plant saponins, where R is comprised of an ethercal ring system

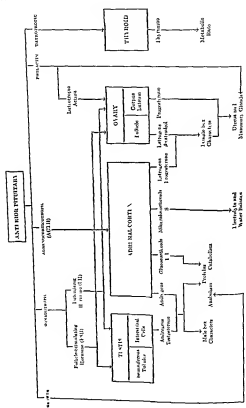
(e) The steroid hormones, where C17 bears a ketonic or hydroxyl group (the androgens and estrogens), or earnes a substituted two-carbon side chain (progesterone and the adrenal cortical steroids)

The steroid hormones all carry phenolie or ketonic oxygen on Ca The summary on pp 751-754 formulates the principal steroid hormones, and illustrates, by means of arrows, the metabolic pathways followed in the hody in the course of their mactivation and excretion. The ovarian follicular hormones—the estrogens—all possess an aromatic or henzenoid ring A, while the hormones of the testis, corpus luteum, and adrenal cortex are all characterized by the presence in ring A of an α-β unsaturated ketonic grouping, the saturation of which leads to practically complete loss of physiological activity. In the course of this reduction, which is a common mechanism of inactivation in vito, carbon atoms numbers 3 and 5 hecome asymmetric, and hence four fully reduced geometrical isomers are possible in each instance. The hydroxyl group resulting on hydrogenation of the ketone may be oriented either cis or trans with respect to the angular methyl group at C10 Respectively, the cis and trans epimers are designated by the suffixes β and α , and conventionally are differentiated in twodimensional formulas by the use of a solid and a dotted line [Compare androsterone (III) and isoandrosterone (IV)] 3(β) Hydroxy-steroids form with digitonin (a saponin from Digitalis) sparingly soluble addition compounds, whereas the enumeric $3(\alpha)$ -compounds do not Likewise the hydrogen atom introduced at Cs on saturation of a 4 5 or 5 6 double hond may he as or trans oriented in relation to the angular methyl group at C10, again, in planar formulation, the solid and dotted lines respectively are used [compare androsterone (III) and etrocholanolone (V)] Because of this isomerism about Cs, there are two saturated parent hydrocarbons of the steroid hormones containing 19 carbon atoms, namely androstane (trans) and etiocholane (cis), and two pertaining to the hormones with 21 carbon atoms, namely pregnane (cis) and allopregnane (trans)

The total synthesis of steroids has been realized through a long sequence of intricate chemical reactions which, however, are not yet applicable commercially Steroid hormones are prepared by degradation of naturally occurring plant and animal steroids Diosgenin, a sapogenin from the Mexican yam, is readily converted to progesterone (XVIII) and deoxycorticostcrone (XIV), and from these to testosterone (II) and estrone (VIII) Much less satisfactorily, the same four hormones may be derived from cholesterol The hile acids, particularly deoxycholic acid, serve as starting material for the part-synthesis of the corticoids hearing an oxygen function at C11, namely cortisone (XV), corticosterone (XVI), and Compound F (XVII) Certain microorganisms possess the uuique and specific property of introducing a hydroxyl group at C11, and they are employed in some commercial procedures for the preparation of cortisone Isotope studies show that in the body all the steroid hormones and cholesterol

can be synthesized from acetate

In general, all storoid hormones are insoluble in water but soluble in fat solvents such as ether, acctone, alcohol, etc., and in vegetable oils, in which medium they are usually dissolved for administration to the body They are relatively mactive per os, and consequently are given intramus-



cularly or subcutaneously. Esters of the hormones, such as the acetate. propionate, and benzoate, are much more valuable theraneutically than the free steroids since their physiological action is more protracted owing to the delayed rate of absorption and utilization. The steroid hormones and their catabolites are generally eliminated in the urine in watersoluble form in conjugation with glucuronic or sulfuric acid.

Formulation of Steroid Hormones and Related Substances. The chemical name is given in each instance, followed by the commonly used name in parentheses. The arrows indicate metabolic pathways followed in the hody.

The Androgens and Urinary 17-Ketosteroids

VII 17(β)-Niethyl-Δ¹-androsten 3-one-17(α)-ol (Methyltestosterone) CuilinO₂

Δ¹-Androsten-3(β)-ol-17-one (Dehydrosoandrosterone) (Transdehydroandrosterone) Cull₁O₁



VIII $\Delta^{1-3.5}$ -Estratriene-3,17(α)-diol (β -Estradiol) C_{11} 11₁₄ O_{8}

XI

Star = Catrapentaen-3-oi-17-one
(Equilenin)

Chili,O

IX Δ^{1,2 t}-Estratriene-3-ol-17-one

ΗΟ X Δ¹ 1.1.-Estratriene-3,16,17-triol

(Estriol) CallaOs

753

trans-3,4(Di-p-hydroxyphenyl)hexene-3

(Diethyl stilbestrol) C18H20O2

Adrenal Cortical Steroids Mineralocorticoids

Δ⁴-Pregnene-3,20-dione-17(α),21-diol (17-Hydroxydeoxycorticosterone) (Compound S) C₁₁H₁₀O₄

Glucocorticoids

XIV Δ^4 -Pregnene-3,20-dione-21-ol
(Deoxycorticosterone) $C_{21}H_{20}O_4$

Δ'-Pregnene-3,11,20-trione-17(α),21-diol (17-lly droxy-11-dehy drocorticosterone) (Cortisone, compound E) C₁₁ll₁₀O₄

Δ⁴-Pregnene-3,20-dione-11,20-diol (Corticosterone, compound B) C₁₁II₂₀O₄

XVII

Δ'-Pregnene-3,20-dione-11(β),17(c),21-triol
(11-Dihydrocortssoe, 17-11ydroxycorticosterone)
(Compound F)

Culls.Ok

XVIII

Δ*-Pregnene-3,20-dione
(Progesterone)

C₁₁H₁₄O₂

X1X
Pregnane-3(α),20(α)-diol
(Pregnanediol)
C₁₁ll₁₄O₂

THE TESTES

Removal of the testes causes atrophy of the seminal vesicles, prostate, and other accessory organs of the male gental system. The effects are most pronounced if the operation is performed before sexual maturity, and are due to withdrawal of androgenic hormones, such as testosterone (II), secreted by the Leydig or interstitial cells. The androgenic hormones will restore the atrophical tissues in cartacted animals, such as rats or meta, and are frequently assayed by their power to cause enlargement of the combining the capon.

Testosterone (II) is the principal male sex hormone isolated from testis tissue. In the course of its metabolism in main it is reduced to androsterone (III), isoandrosterone (IV) and etiocholanolone (V), all of which are exerted in the uriue (see 17-ketosteroids, p. 759). Methyltestosterone (VII), prepared artificially, but not known to occur in nature, finds wide therapeutic application because it is highly active per os

The International Standard male hormone unit is the activity equivalent of 0.1 mg of pure androsterone. By comparison of comb growth in the capon, testosteroue is about six times, and isoludrosterone oueseventh, as potent as androsterone, while etocholanolone is practically inert. Administered parenterally, methyltestosterone and testosterone show equal activity, orally methyltestosterone is two to four times less active than parenterally, but about 20 times more potent than testosterone given by mouth

THE OVARIES

The development of the accessory sev organs in the female is not so dependent on hormonal stimuli as in the male, nevertheless, extripation of the ovaries causes atrophic change in the uterus, vagina, mammary glands, etc., and in the pluntage of poultry. In most mammals, mating occurs only during periods of heat or estrus, when the concentration of estrogenic hormones produced by the ovary and in particular by the Graafian follicles becomes maximal. In rodents estrus is accompanied by the appearance of coruffed cells in the lumen of the vagina, from the walls of which they have desquamated. This reaction follows the administration of estrogenic substances to castrated animals, and is the hasis of hiological assay.

After ovulation in mammals, the ruptured Graafian follicle is transformed into the corpus luteum, which produces the hormone progesterone (XVIII) whose chief function is to prepare the mucous membrane of the uterus for the implantation of the embryo Progesterone is assayed by its power to cause a glandular "progestational" prohferation of this membrane in immature or castrate rabbits previously sensitized with estrogens Gestation, in its early stages is interrupted by destruction of the corpora lutea, when implantation and pregnancy occur, the active hie of the corpora lutea is prolonged In the later stages of human pregnancy, estrogens and progesterone are produced by the placenta, they act together in promoting mammary development

The principal estrogens are β -estradiol (VIII), estrone (IX), and estrol (Δ), their relative activities are roughly 10-1-1 or less, but wide fluctuation in these proportions is observed by the different methods of testing Only the first two are known to occur in liquor folliculi, where β -estradiol accounts for 90 per cent of the physiological activity, and hence is regarded as the chief or arian follicular hormone β -Estradiol and estrone are interconvertible in the animal organism, for the administration of either leads to the exerction of the other. Estrol has been isolated only from human pregnancy urine ind from human pheenta, and that organ is regarded is the main site of production. In gestation (Fig. 202), the estrogen output rises sharply with the growth of the placenta to a pre-

partum level of 12 to 10 mg per diem, estrol constitutes the chief catabolite (about 90 per cent of the total), but estrone is exercted in appreciable amount together with traces of β-tstradio! The conjugate of estrol in human pregnancy urine has been isolated as the 16- or 17-monoglucuronide. Throughout the normal menstrual cycle, the urine estrogen contents low, of the order of 0.08 mg per diem at the maximum. Biological assisticted in the period of flow. While the chemical nature of the estrogens exercted in the nonpregnant state has not been established by isolation, partition experiments indicate that all three are climinated.

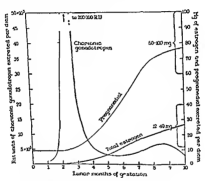


Fig 202 Variation in Universit Output of Sex Hormones and Metabolites with Period of Gertation

Throughout gestation in the mare, estrone represents the most abundant exerctory product Estrol is not formed in this species, but considerable amounts of estrogens unsaturated in ring B as well as in ring 4 are eliminated, equilent (XI) is the most abundant

Many synthetic derivatives of stilbene possess marked estrogenic potency, although they are nonsteroidal and do not occur in nature Important among these is diethyl stilbestrol (XII) which, like other mem bers of the group, is much more active orally than the free natural estrogens. The water-soluble conjugated estrogens of human and equine pregnancy urine (principally estriol glucuronide and estrone sulfate respectively) are also effective by mouth.

The International Unit is defined as the activity contained in 0.1 μ g of estrone. As the different estrogens cannot properly be compared because of variations in the time and period of action, a subsidiary standard

exists for β -estradiol-3 monohenzoate which is the activity equivalent of 0.1 μ g of that substance

Progesterone is produced by the corpus luteum of the ovary during the postovulatory phase of the menstrual cycle, and in much larger amounts throughout gestation by both the corpus luteum and the placenta Traces are elaborated by the adreoal cortex to both seves. The chief metabolite of progesterone (AVIII) is the loactive pregnancdiol (XIX), which is excreted in the urine in conjugation with glucuronic acid. Estimation of the sodium pregnanedial glucuronidate content of urine thus provides a useful index of luteal function From 40 to 55 mg of pregnauediol are eliminated during the normal menstrual cycle, the excretion beginning probably one to two days after ovulation and ceasing one to three days before the onset of uterine bleeding A significant decrease in total output, or shortening of the period of excretion indicates that progestation is not sufficiently well developed or is of too brief duration to support implantation From fertilization to the end of the ovariao phase of pregnancy (ca three to four lunar months), urme pregnanediol remains at the maximum level associated with the luteal phase of the cycle (ca 5 to 10 mg per diem), and then rapidly rises with the growth of the placenta to a daily output at parturation of about 60 to 100 mg (Fig 202) A significant fall during the first third of gestation indicates corpus luteum deficiency or failure of the placenta to develop at the normal rate, either of which threatens abortion. A residual pregnauediol titer of the order of 0.2 mg per diem remains in the urine of men and of women in the follicular phase of the cycle, presumably this is of adrenal cortical origin. The loternational Unit is defined as the activity contained in 1 mg of the pure crystalline progesterone

THE ADRENAL CORTEX

The adrenal glands consist of two distinct tissues in close apposition the medulla, producing epinephrine (adrenaline) and norepinephrine (noradrenaline) and derived from the sympathetic nervous system (see p 766), and the much larger surrounding cortex, embryologically related to the gonads Extirpation or destruction of the cortical tissue is fatal Before death, the body loses sodium salts in the urine and accumulates potassium, so that the volume of extracellular fluid is reduced the blood becomes viscous the circulation sluggish, and renal function seriously impaired, treatment with diets low in potassium and enriched with sodium salts is helpful. Another group of symptoms includes diminished hreakdown of tissue proteins to form glycogenie amino acids, so that glycogen stores decrease rapidly on fasting hypoglycemia develops and resistance is greatly lessened to extremes of heat and cold, infections, toxic drugs, traima, fatigue, and stress of all kinds. Addison's discase, in which the cortical tissue is gradually destroyed, is characterized by asthema emaciation, low blood pressure, pigmentation of the skin, and often hypoglycemia, although here (as in the adrenalectomized dog) the disturb nice of electrolyte metabolism is usually the greatest threat to surva al

About 30 steroids have been isolated in crystalling form from cortical

extracts Of these 30 steroids, many are physiologically quite mert, others, such as traces of estrone (IX) and progesterone (XVIII) have heen discussed above, several, such as androstenedione, adrenosterone, and 17 hydroxyprogesterone, have definite androgeme activity, but the most interesting are six which are active in correcting the specific effects of adrenalectomy and may therefore be called corticoids (Selye) All six are derivatives of A4-pregnent 3,20-dione-21-ol The adrenal contains enzyme systems which promote the introduction of oxygen at positions 11, 17, and 21 of the steroid skeleton, for example, adrenal tissue, in perfusion or incubation experiments, will readily convert progesterone (XVIII) to compounds I (XVII), E (XV) and B (XVI) Deoxycorticosterone (XIV) is present in very small amounts, if at all, but as it is readily available commercially, it has been widely used in medicine, especially as its 21acctate (DCA), it is highly potent in correcting the imbalance in the metabolism of sodium and potassium and hence in maintaining life in the adrenal ctomized dog or in patients with Addison's disease, and may be called a mineralocorticoid On the other hand, cortisone (17-hydroxy-11-dehydrocorticosterone, Kendall's "compound L") (XV), and hydrocortisons (17-hydroxycorticosterone, Kendall's "compound F") (XVII) may be called glucocorticoids, they have slight and variable action on sodium and potassium balance but profound influence on protein and carbohydrate metabolism, on which their medical uses depend Corticosterone (XVI) and 11-dehydrocorticosterone possess both glucocorticoid and mineralocorticoid activity (oxygen at position 11 is essential for the former), but not in any high degree It must be emphasized that cortical extracts contain mineralocorticoids with life maintaining activity These compounds (the "amorphous fraction") bave not yet heen crystallized or identified and are much more water soluble than the glucocorticoids Biological assay procedures for glucocorticoids are based on their power to increase resistance to cold in adrenalectomized rats or to restore liver glycogen levels. Mineralocorticoids are assayed by their life-maintaining power or more specifically by their effect on the metabolism of sodium and potassium

The structure and secretory activity of the adrenal cortex is controlled by the adrenocorticotropic hormone (ACTH) of the anterior pituitary After hypophysectomy, the cortex atrophies, especially the inner layers and low resistance to stress and impaired glyeogen storage indicate that glucocorticoids are not being produced. The injection of a single dose of ACTH, in the rat, leads to rapid depletion of the cholesterol and ascorbic acid of the cortex (this is the basis of the Sayers assay method) with some temporary increase in cortical volume and various evidences of increased manufacture and discharge of corticoids, principally hydroxycorticosterone (compound b) The application of suitable stress, e.g., brief exposure to cold or the injection of a sublethal dose of epinephrine, will produce the same results by causing the pituitary to discharge ACTH In human subjects the injection or discharge of ACPH or of glucocorticoids is evidenced by (a) a fall in the numbers of circulating eosinophils and lymphocytes (b) an increased exerction of nitrogenous waste products, notably une acid, and (c) increased urinary excretion of corticoids as determined by chemical or biological assay (but representing only a small fraction of the corticoids actually secreted or injected) and of 17-ketosteroids ACTH and compounds E and F have proved dramatically successful in relieving the symptoms of rheumatoid artbritis and many other diseases, the disappearance of pain and the development of a sense of well being are particularly remarkable. Prolonged treatment with small daily doses of cortisone appears to be both possible and effective However excessive doses administered over a long period of time may induce symptoms of Cushing's syndrome, with tendencies toward obesity. hypertension, osteoporosis, and psychic disturbances. In patients disposed to diabetes, symptoms of this disease may be aggravated. Large doses are contraindicated in tuberculosis, gastric ulcer, and diabetes Cushing's syndrome is thus attributed to overproduction of glucocorticoids The concept of a eausal relation between overproduction of mineralocorticoids and the collagen diseases (rheumatoid arthritis, periarteritis nodosa, etc.), attributed to Selve, has been questioned by many investigators and is the subject of active study. Overproduction of adrenal androgens, with high 17 ketosteroid excretion, leads to the adrenogenital syndrome (virilism, hirsulism, or precocious masculinization) with cnhanced protein synthesis and museularity

17-KETOSTEROIDS

Human urines contain steroids earrying ketonic oxygen at C₁₇, some of which are phenolic while others are ueutral. The latter are commonly called "17 ketosteroids." The principal members of this group are androsterone (III), etocholauoloue (V), isoaudrosterone (IV), and dehydroisoandrosterone (VI), the first two of these arise in part from the testis, while all represent the exerction products of some of the steroids of the adrenal cortex. Accordingly, measurement of 17-ketosteroid output provides a biochemical index of testicular and adrenocotical activity.

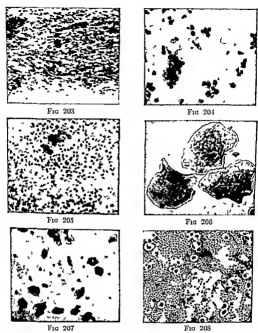
The normal 17-ketosteroid excretion of men between the ages of 20 and 40 years averages about 15 mg per duem normal women in the same age group excrete approximately 10 mg per duem. In children under 8 years of age, less than 1 mg per duem is eliminated, but from this age on there is a gradual increase to adult values. Likewise in old age a significant diminution is observed. As gonadectomy in the male decreases the average output from 15 to 10 mg, and is without effect in the female, it is concluded that approximately 10 mg are derived from the adrenal cortex and 5 mg from the testis. The human ovary is not a source of neutral 17-ketosteroid.

Disorders of the testis, adrenal cortex, and anterior pituitary may profoundly after 17-ketosteroid exerction. In eunuchoidism values from normal to that of the surgical castrate (10 mg) are reported whereas in the rare cases of masculinizing tumors of the interstital cells of the testis the output may reach 800 mg per diem. In Addison's discuse in the male, exerction fells to 1.2 to 6.4 mg, which represents the testicular output, while in the female practically no 17-ketosteroids are produced. In those cases of Cushing's syndrome not associated with carcinoma of the adrenal cortex, normal or only slightly elevated 17-ketosteroid values are ob-

served (10 to 36 mg), but when caremoma of the cortex complicates the condition a much higher exerction is usually encountered (40 to 28 mg) Similarly, in the adrenogenital syndrome, simple hyperplasia of the cortex leads to only a moderately high 17-ketosteroid output (up to 100 mg approximately), whereas caremoma generally gives rise to a more marked increase (ca 100 to 250 mg). In both instances, caremoma may be distinguishable from hyperplasia by the higher exerction, and also by the increased proportion of the $3(\beta)$ -hydroxy-17-ketosteroids (chiefly dehydroisonadrosterone) to the $3(\alpha)$ -hydroxy-17-ketosteroids (chiefly a ratio is 19, whereas in caremoma it may rise to about 11 In pan hy popituitarism, a general underproduction of all anterior lobe hormones the 17 ketosteroid excretion is low (0 to 3 mg.)

EXPERIMENTS ON STEROID HORMONES

1 Partition and Estimation of the Estrogens of Urine Principle The determination comprises (a) hydrolysis of the water-soluble conjugates in urine (b) extraction of the skirod more ties and it en separation by partition between solicing and (c) estimation of the estrogen content of the various fractions either colorimetrically (see Txp 3) or by hological assay (see 1xp 4) Colorimetric methods are unsatisfactory when the estrogen output is less than 15 mg per liter approximately as in n irmal urine and throughout the first four months of gestation



Figs 203-208 Vaginal Smears of Artificially Induced Estrus in Spated Rat

Fig 203 Diestrous smear leukocytes in stringy mass (× 40)

Fig 204 Proestrous smear chefly nucleated epithelial cells with an occasional leukocyte (× 40) Present 35 to 40 hours after first injection.

Fig. 205 Estrous smear nonnucleated cornified epithelial scales (× 40) This type usually appears within 48 hours after the first injection and is a certain criterion of the positive action of an extract

Fig 206 Flat cormfied elements of the estrous smear stage (× 250) Losin stans these cells a brilliant red. Uthough the site of the former nucleus is apparent all hasophile staning reaction has been lost

Fig 207 Early stage of leukocytic infiltration (metestrum) (× 40) Few nucle ited cpithchal cells have appeared as yet

Fig. 208 Late stage of the metestrum (× 40) Fuormous numbers of lunkocytes some coronfied scales (in center of field), and many nucleated epithelial cells

From Allen, Douy et al. Am, J. inat. 34 163 (1924-1925) Courtes) Cameron Recent. Advances in Endocrinology 5th ed. Philadelphia, The Blakuston Company. 1945.

washed successively with 12 ml. of 45 per cent II₂SO₁, twice with 25 ml. of 9 per cent N₂CO₁, and twice with 25 ml. of water, and evaporated to dryness (Fraction OD).

Both Fractions T and OD are purified sufficiently that they may be assayed colorimetrically (see Exp. 3). When a separate estimate of the estradiol and estrone content is required, Fraction OD is divided into ketonic (estrone) and nonketonic (estradioi) fractions by treatment with Girard's reagent? prior to assay.

- Isolation of Crystalline Estriol from Human Pregnancy Urine. Fraction
 T (Exp. 1) from late pregnancy urine readily yields reasonably pure estriol
 on crystallization from benzene. For this purpose 500 ml. or more of urine
 may be taken, ail quantities of reagents and solvents being increased proportionately.
- 3. Kober's Color Reaction for Estrogens. The sample to be tested is evaporated to dry ness in a test tube, to which is added 2 mi, of an equimolecular mixture of sulfurlicand (o and p) phenylsulfonic acids. The test tube is heated for ib minutes at 100° C. and at once cooled in ice. A red color gradually develops.
- 4. Biological Test for Estrogens. Young adult female rats or mice are ovariectomized. If the operation is complete, within a week the vaginal smear will contain only leukocytes. The smear may be obtained by washing out the vagina with a drop of saline solution, using a small homemade pipet with a rubber bulh, or by introducing a small pledget of moist cotton wool on a loop of platinum wire, and in either case spreading the fluid on a clean microscope silde. It is not necessary to fix or stain the cells. Inject the animals with three 0.5 ml. doses of a solution of estrin in oil, or in 10 per cent alcohol made slightly aikailne, preferably allowing at least 6 bours to elapse between injections. Examine the vaginal smear on the second, third, and fourth days. The leukocytes will almost disappear, and the smear should come to consist of horny, scale-like squamous cells, and some epithelial cells still retaining their nuciel. The test can be made quantitative only if large numbers of animals are used, and their sensitivity to crystalline estrogens must be established. The response is affected by the strain of the animals, by the degree of subdivision of the dose and the soivent used, by the number of examinations made, and by many other factors. See Figs. 203-208.

pressure. The residue is taken up in 60 ml. of 0.1 N NaO11, again extracted 3 times with butanol (60, 20, and 10 ml.), and the combined butanol extracts are washed twice with 5 ml. of water and evaporated to dryness under reduced pressure. The residue is dissolved in exactly 5 ml, of water and transferred to a 125-ml, flask with several small portions of acetone. Volume is made to 100 ml. with acetone, and, after standing overnight in the refrigerator, pregnanediol glucuronidate precipitates. Most of the supernatant acetone is removed by suction without disturbing the precipitate. The remaining mixture is transferred to a 50-ml, centrifuge tube and centrifuged for 10 minutes. The residual acetone is carefully decanted and the crude ester is dissolved in water for purification by reprecipitation. The amount of water used is dependent upon the weight; 2 ml. for quantities less than 5 mg.; 3 ml. for 5 to 10 mg.; and 5 ml. for more than 10 mg. The water is added to the original flash to dissolve adhering material and then transferred to the 50-ml, centrifuge tube containing the precipitate. The tube Is warmed and the contents are filtered back with suction into the original flask. The tube is rinsed twice with small amounts of acetone which are filtered into the flask, and the volume is made to 100 ml, with acetone. After 12 hours in the refrigerator. the acetone is removed as before by auction and centrifugation. The precipitate is dissolved in 10 ml, of hot 95 per cent ethanol and filtered with suction into a tared beaker (30 ml.). The flask, centrifuge tube, and filter paper are washed twice with hot ethanol. After evaporation of the solution to dryness on the bath, the weight of the precipitate is determined.

CALCULATION. The percentage recovery of the ester depends upon the amount present and the volume of water used in the second precipitation, accordingly a correction factor is applied (see the table below) Sodium pregonoediol glucurouidate contains 61 7 per cost pregnancediol

Mg pregoacediol excreted per diem = \frac{\text{Wt of ppt \times 0 617 \times 24 hr volume \times 100}}{\text{Per cect recovery \times volume extracted}}

The melting point of the final precipitate should always be coofirmed, sodium pregnanediol glucuronidate melts at 273° C, with evolution of gas.

Weight of	Per cent recovery				
precipitate in mg		2 ml, H=0	3 ml H ₂ O	5 ml H:0	
_	2	0-60*			
	3-4	50-67*		-	
	5-8	70-75*	79		
-	9-10		82	75	
	11-12		85	78	
	13-15			81	
Ξ	16-18			83	
	19-25			85	

^{*} Approximate values.

- 6. Preparation of Adrenal Cortex Extract: Method of Cartland and Kuizenga. Minced whole beef adrenals are thoroughly extracted with 99 per cent acetone (2.5 liters per kg.) and again with 80 per cent acetone. The extracts are combined, filtered, and concentrated under reduced pressure (helow 45° C.) till the acetone is removed. The aqueous residue is twice extracted with petroleum ether, which removes inert fats and is discarded, and then twice extracted with ethylene dichloride (400 ml. per kg. gland each time), which extracts the hormone but not adrenaline nor phospholipides. The ethylene dichloride soluble fraction is chilled to -15° C., and ice and precipitated solids are filtered off. The ethylene dichloride is removed under reduced pressure and the residue dissolved in ethyl alcohol. This is mixed with an equal volume of petroleum ether; water is then added to make the alcohol 90 per cent; this causes some of the petroleum ether to separate as an upper layer, which is removed in a separatory funnel and discarded. Two further discarded petroleum ether fractions are obtained by diluting the alcohol further to 80 and finally 70 per cent. The alcohol is now removed under reduced pressure, below 45° C., and the resulting aqueous colloidal solution sultably diluted to give a final preparation of volume 1 ml. per 40 g. gland, with NaCl added to make 0.9 per cent, and containing 10 per cent ethyl alcohol as preservative. This final solution is filtered and sterilized by Berkefeld filtration, and preserved in sterile ampoules in the refrigerator. (For details of this method, see the original paper.)
 - 7. Chemical Assays of Corticoids: Principles. Several colonmente methods have been described all of which depend upon the Car-Cui kctol grouping in the side sham but are independent of the functional groups in the ring structure. These are based upon (a) reduction of phosphomolyhdic and to molybdenum blue, (b) reduction of cupric onto ecuprous ion, and (c) evidative cleavage with penodic sed of carbon atom 21, which is released as formaldchyde and estimated with chromotropic and The methods are applied to the full 21-thour output of urns and lead to itter of "reducing steroids" and "formaldchydogeme steroids" All the methods melide in the estimates the active corticoids, plus much larger quantities of inactive methods in the estimates the active corticoids, plus much larger quantities of inactive methods by the respond to these reactions but are physiologically mert. Consequently the values determined by chemical assay (about 1 to 2 mg per day in normal indrividuals) are considerably higher than the glucocorticoid tier (See footnote 1, p. 760).
 - 8. Biological Assay of Clucocorricoids and Determination of Urinary Cortin by Liier Glycogen Deposition: Method of Venning, Kazmin and Bell. The muthod is applicable to crystalline corticoids or to unne extracts. The reference standard is cortisone (N1), the biological activity of 1 pg of which is defined as one glycogene unit in the assay of unknown extracts.

Procedure. For urines containing a normal or low titer, a 48-hour specimen is required. The urine is adjusted to pli 1 with HCl, and extracted 3-4 times with ethylene dichloride or chloroform. Any emulsions encountered can be broken by centrifuging. The clear extracts are evaporated almost to dryness under reduced pressure (water bath temperature not over 55° C.). The residue is taken up in 30 ml. of chloroform, and the solution is extracted 3 times with

¹ Cartland and Kuizenga J Biol Chem 116, 57 (1936) See also Kuizenga, Wick, Ingle-Nelson and Cartland J Biol Chem 147 561 (1943)
² Venning Kazimi and Bell Endoctrinology 33, 79 (1946)

5 ml. of cold 0.1 N NaOH and 3 times with 5 ml. of water. These washings are back-extracted with chloroform, the chloroform extracts combined, and evaporated to 1-2 ml. for transfer to a test tube. Finally they are evaporated to dryness under a stream of nitrogen. The dry residue may be stored until ready for assay.

For the higassay male white mice weighing 20-25 g, are used. Two days hefore adrenalectomy the mice are taken off a stock dlet of "Purina" chow and put on the McCollum lactation dlet, which contains 26 per cent protein and 52 per cent carbohydrate. Removal of the adrenals is effected by the usual lumbar route. Following the operation, the mice are kept in a constant. temperature room or box at 76° F., and maintained on the McCollum dier supplemented with NaCl (0.9 per cent) and clucose (5 per cent) in the drinking water for the first postoperative day. Glucose is withdrawn thereafter: salt is retained in the dlet throughout the test. Food Is removed at 5 P.M. of the third postoperative day, and the mice starved until the following morning. Drinking water is removed on the fourth postonerative day, and beginning at 9:15 A.M. a total of 7 Injections are given at 9:15, 10:00, 10:45, 11:30 A.M., 12:30, 1:30, and 2:30 P.M. The material to be tested is taken into solution containing 5 per cent glucose and 10 per cent alcohol. At each injection 0.20 ml. is given subcutaneously, so that each mouse receives a total of 1.4 ml. of extract containing 70 mg, of glucose. At 3:30 p.m. the mica are weighed and anesthetized with Sodium Amytal (0.2 ml, of 1.8 per cent solution). The livers are quickly removed and plunged into 4 ml, of hot 30 per cent KOII contained in a 15-ml. graduated centrifuge tuhe. The tubes are heated in a boiling water bath and frequently shaken, until all the tissue is in solution. The glycogen is precipitated by the addition of 1.2 volume of 95 per cent alcohol. The tubes are reheated until the mixture just begins to boll, cooled in an ice bath, and centrifuged. The supernatant liquid is poured off, and the tubes are allowed to drain. The sides of the tubes are washed down with 0.5 ml. of alcohol and again allowed to drain. Final traces of alcohol are expelled by heating tha tubes for a few minutes in the hot water bath.

The glycogen is hydrolyzed as follows: After addition of 5 ml. of 1 N H. SO₄, the tubes are placed in an autoclave and heated for 15 mlnutes at 15 lb. pressure. The glucose is determined by the method of Good et al. Nor colorimetrically by the method of Nelson's (see p. 573). The glycogen is expressed in terms of mg. of liver glucose per 100 e. of mouse body welcht.

Six to eight mice must be used for each assay. For normal male urine the equivalent of 6 hours of urme is administered to each mouse, whereas for normal female urine expected to bo low in glycogenic activity, the equivalent of an 8-hour allouot of urine is given each animal.

Values found by this method range from 0.04 to 0.085 mg. per 24 hours for normal men, and from 0.025 to 0.065 for women, calculated as cortisone.

9. Determination of the Neutral 17-Ketosteroids of Urine: Principle. The steroids of urine are first set free from their water-soluble conjugates by each by drolly sis and extracted with other. After removal of pheodic and eachie impurities, the 17-ketosteroid content of the neutral residue is ascertained by quantitative application of the Zimmerman reaction. This comprises coupling of the reactive 17-ketone group

Cited by Agate and Zwemer, Am J. Physiol, 111, 1 (1935).
 Good, Kramer, and Somogyi J. Biol. Chem., 100, 485 (1933).

¹¹ Nelson J. Biol Chem , 153, 375 (1944).

¹² Zimmermann; Z. physiol, Chem., 233, 257 (1935).

of the compounds with m-dimitrobenzene to form colored complexes, the intensity of which in the green region of the spectrum (a20 ma) is measured in the photoelectric colorimeter and compared with the color developed by known amounts of a pure crystalline I7 ketosteroid

Nonspecific chromogens of urme cause some interference, but as these substances are mainly nonketonic and absorb light m the volet (420 m) as well as the green (520 m) region of the spectrum, their miluence may be eliminated either by carrying out the color determination on the neutral ketonic fraction of urme, or by the application of a correction factor, which, in effect, subtracts the color component due to extraneous ethoronogens

Procedure 12 A 24 hour specimen of urine is collected and measured. To 2 100 mi aliquot, 15 mi of concentrated HCl are added The mixture is re fluxed for exactly 15 minutes, and then immediately chilled and extracted 4 times with 30 mi portions of freshly redistilled ether The combined ethers are washed 4 times with 2N NaOll (15 ml portions) and then twice with water On evaporation of the other, the neutral residue is dissolved in 10 ml ketone free ethanol in the development and measurement of the color, matched colorimeter tubes must be used To the experimental tube are added, in order and with mixing after the addition of each, 0 2 ml of the urine extract, 0 2 ml of 2 percent pure m dinitrobenzene (prepared fresh) in ethanol, and 0 2 mi of 5N aqueous KOH (purest electrolytic grade) A control tube 18 made up in the same way containing 0 2 ml of ethanoi in place of the urine extract Both tubes are incubated in the dark at 25 ± 0 5° C for exactly 105 minutes Then 15 ml of 80 per cent ethanol are added to each, and, after mixing the color intensity in each is measured in a suitable photoelectric colorsmeter To correct for the slight color developed by the readents alone, the control tube is first inserted in the instrument with the 520 mm fitter in position, and the instrument adjusted to 0 density (100 per cent transmit tance) The control is now replaced by the experimental tube, the reading noted and referred to the calibration curve pertaining to pure androsterone to give the equivalent number of mg of this substance in the sample

CALCULATION

Mg 17 kctosteroids exercted per diem = Mg observed × ml 24-hour urine volume

For the application of factors correcting for interfering chromogens, readings are taken at 420 mm also. For further details and modifications see footnote 1, p. 760 when differential estimates of the 3(a) and 3(b) hydroxy 17 ketosteroids are desired their separation is effected with digitionin more to assay

NI FROGEN-CONTAINING HORMONES

ADRENAL MEDULLA

This organ which gives a specific staining reaction with dichromates ('chromaffin cells'), contains the active introgenous bases epinephrine or adrenaline ('N) and norepinephrine (noradrenaline arternal) (N'1). Both may also be prepared synthetically, the latter appears to be more abundant in the circulating blood

¹² Method used in the McG II University Clime.

The natural forms are levorotatory and are many times more active than their optical isomers. They are basic substances, insoluble in weak alkalies and in organic solvents, and form water-soluble crystallizable salts with various acids They are readily destroyed by oudation in alkaline solutions, eventually giving rise to amorphous, dark-colored melanins Being attacked both by "amine-oxidase" and by the cytochrome system they are quickly destroyed in the body, so that the effect of an intravenous injection rapidly passes off, they are slowly and incompletely absorbed from subcutaneous or intramuscular injection sites, where they are ant to produce abscesses, they are mactive orally though somewhat toxic The physiological reaction to these hormones is, in almost all organs, the same as the response to stimulation of the sympathetic nerve supply the blood pressure rises because of arteriolar constriction, the heart rate increases if the depressor reflex can be excluded, peristals is inhibited, the pupils dilate, the brouchial muscles relax, there is outpouring of ACTH and probably other anterior pituitary hormones, blood sugar and blood lactic acid increase at the expense of the glyeogen stores, and the metabolic rate is temporarily increased. Norepinephrine shares the stimulatory rather than the inhibitory properties of the better-known coinephrine Discharge of these hormones occurs only when stimuli reach the adrenal medulia by way of the splanchme nerve This happens rapidly when the body is exposed to emergencies such as cold or shock or violent emotion or pain

EXPERIMENTS ON THE ADRENAL MEDULLA

1. Preportion of Epinephrine To each 100 g of ground fresh adrenal tissue add 50 ml of a 3 per cent solution of acetic acid in 95 per cent ethyl alcohol Allow to stand for 12 to 24 hours and remove the liquid by straining. Restract the residue twice for 6 hours, each time with the same volume of 3 per cent acetic acid in 60 per cent ethyl alcohol as before After filtering, concentrate the extract in a flask under diminished pressure to about one-tenth of the original volume. The material which has precipitated should be removed by filtration and the filtrate concentrated under diminished pressure to about 2 to 3 ml per 100 g of tissue extracted. Transfer the solution to a test tube and add enough strong ammonium hydroxide to leave a strong odor of ammonia. Stopper and set aside in a cool place for several hours. Remove the precipitate by filtration and wash first with lee-cold water which has been

boiled then with cold alcohol and finally with ether Dissolve the precipitate in 10 ml of 10 per cent hydrochloric acid, reprecipitate, and treat the precipitate as before

- 2 Properties of Epinephrine Dissoive 0 1 g of epinephrine in about 10 ml of 0 1 per cent HCl and dilute to 100 ml
- a VULPIAN REACTION Add a few drops of FeCI, solution A green color will be produced which is a typical catechol reaction
- b Fwins Reaction Add to I ml of the solution an equal volume of 0.2 per cent potassium persulfate a reddish color is produced which is specific and may be quantitatively determined "
- c FILOMESCINCE. A faintly alkaline solution of epinephrine even if extremely dilute shows a bright apple green fluorescence for some minutes when irradiated in a quartz ressel with ultraviolet light from a mercury vapor jamp

THE PANCREAS

Removal of the pancreas in eats and dogs produces symptoms similar to diabetes mellitus in man. Blood sugar increases greatly and glucosuria



Fig 200 Crystalline Zinc Insulin Courtesy Dr D A. Scott

occurs the glycogen stores of hier and muscle are depleted the blood is charged with fat and acctone bodies (acctoacetic acid \$B\$-hydroxy butyro acid and acctone) accumulate in the blood and are exerted in the urne to produce acidosis coma and death usually within three weeks These symptoms do not follow when the digestive pancreatic juice is prevented from reaching the intestine or even when the acinar cells which secrete this juice digenerate they are rather due to removal of solid clumps of cells the islets of Langerhans which have no connection with the ducts but form a hormone insulin which is discharged into the blood and regulates carbohy drate metabolism. Diabetic symptoms appear when the late cells are damaged by administration of alloxan or in some species by prolonged administration of certain anterior pituitary extracts which produce hypergly cenns.

The first useful pancreatic extracts were prepared by extracting the pancreas with acid alcohol and increasing the concentration of alcohol in the filtered extracts until insulin was precipitated Refinements of the method led to the isolation of insulin in crystalline form as a zine salt (hig 209) Insulin is a protein somewhat of the albumin type but unusually soluble in moderately dilute alcohol and acctione It is destroyed by digestive enzymes and must be proteited from pancreatic trypsin during extract in lience also it is miffective orally. Insoluble compounds of in thin with banc proteins (protamines or globin) and zine are used therapeutically since they are slo ity absorbed from the tissues and injections need not be given so frequently. In normal animals insulin lowers

¹⁴ Barker Lastland and Evers Buchem J 26 2129 (1932)

the blood sugar, thus eventually producing convulsions unless counteracted by the administration of glucose, or substance yielding glucose. The unit was at one time defined as one-third of the amount which will, in five hours, lower the blood sugar of a fasting right to the convulsive level (45 mg per 100 ml), but assay is now conducted by comparing the activity of an unknown sample with that of crystalline insulin, either in lowering the blood sugar of rahhits or in indusing convulsions in mice crystalline insulin is reckoned it 22 units per mg

Some consider that insulin is an essential catalyst in the hiological oxidation of carbobydrate, for example, that it favors the action of the enzyme hevolutase in forming glucose-6-phosphate as a first step, others believe that insulin checks the new formation of carboby drate from fat (at least from glycerol) and protein, which in the absence of insulin is supposed to flood the organism with sugar, others again hold that lack of insulin inhibits reactions wherehy a large proportion of detary carboby drate is converted into fat. The consequences of pancreatectomy are less marked in animals from which the anterior pituitary, or the adrenal cortices, are removed, they are in any case less marked in species other than the cat and dog

Panereaux extracts, including most insulin preparations, contain a hyperglycenic factor (HGF) which promotes the breakdown of liver elycogen, it has been suggested that this too is a hormone, produced by the α cells of the islets, insulin almost certainly arising in the β -cells

EXPERIMENTS ON THE PANCREAS

- 1 Preparation of Insulin Method of Jephcott " To 1 kg or more of fresh, finely minced beef pancreas, add 4 volumes of extraction liquid, consisting of 750 ml of ethyl alcohol, 250 ml of distilled water, and 15 ml of concen trated hydrochloric acid Shake or stir for 2 hours at 37° C and filter through a double layer of cheesecloth, extract the residues as before Combine the filtrates and add concentrated ammonia till alkaline to litmus, centrifuge, and discard the precipitate (finsulin may be precipitated quantitatively by adding 11/2 volumes of absolute ethyl aicohol and 21/2 volumes of ether and allowing to stand in the cold) A purer preparation is obtainable as follows After centrifuging, drive off the alcohol with a blast of air at 37° C and add 40 g of ammonium sulfate per 100 ml The precipitate which rises to the top contains the jusulin, it is ground with 70 per cent alcohol and filtered To the filtrate add an equal volume of 95 per cent alcohol, and discard any precipitate which forms, now add 8 volumes of 95 per cent alcohol to precipitate the Insuila, which is filtered off and dried, and dissolved in water (1 mi per 10 g of pancreas) containing 0 1 per cent tricresol
 - 2 Effect of Insulin on Blood Sugar Take a 1 ml sample of blood for a sugar determination from the marginal ear vein of a 2 kg rabbit which has been starved for 24 hours. Then inject subcutaneously 5 to 10 units of insulin Observe the rabbit carefully for symptoms of hypoglycemia, such as hyperiritability, palpitation of the heart, convulsions, and coma. Another sample of blood should be drawn as soon as distress is evident and a third when the rabbit is in convulsions. To relieve the induced hypoglycemia, inject 16 ml

¹ Jepl cott Tr Roy Soc Ca ala sec 5 25 15 (1931)

of 10 per cent glucose solution intravenously. After recovery, take another blood sample for a sugar determination Compare the blood sugar values, it is also possible to save the animal by injecting 0.5 ml of 1.1000 epinephrine (adrenaline) subcutaneously after the first consulsions, or convulsions may be averted by feeding glucose upon a leaf of lettuce, or by giving glucose by stomach tube or intraperitoneally.

THE THYROID

The thyroid gland consists of a framework of connective tissue enclosing numerous vesicles lined with epithchum, flattened in the resting state and cubical or columnar (almost obliterating the colloid-filled cavit) of the vestele) when highly active Marked enlargement of the gland



FIG 210 EFFECT OF THIBOXINE ON MIXEDLMA The time interval between pictures is three weeks. The total amount of thyroxine used was less than 20 mg. From Lendall Ind. Eng Chem 17, 525 (1924)

is known as gotter it may be associated with normal or subnormal activity, in which case it is related to a deficient intake of iodine and is chiefly found in inland regions, for example around the Great Lakes, where the soil, water, and vegetation are iodine-poor It may also be associated with increased thyroid activity (Graves's disease, exophthalmic goiter) The thyroid is far richer in iodine than any other tissue The iodine is chiefly built into the characteristic protein thyroglobulin, which makes up a large part of the colloid in the vesicles and may be isolated in relatively pure form 16 Thyroglobulin on bydrolysis yields thyroxine

[&]quot; Heidelberger and Palmer J Bad Chem 181 473 (1933)

or β -[3 5-duodo-4-(3' 5'-duodo-4'-hydruxy-phenoxy)-phenyl]- α -ammopropionic and This is usually obtained in the racenic form, but is originally levorotatory. Truodothyronine, containing one iodine atom less,
is also present and highly active. The administration of thyroxine or of
substances containing it produces only in part the picture of Graves's
disease high metabolic rate, increased pulmonary citilation and circulation rate tending to overwork the heart, extreme nervous restlessiess. and

sometimes protrusion of the eveballs (cyophthalmos) Deficient thyroid ac tivity produces mysedema (see Fig. 210) with low basal metabolism, mental and physical sluggishness, and forma tion of a curious puffy tissue under the skin The extreme form, in which severe deficiency dates from infancy, is known as cretinism (see Fig 211), and is marked by stunting, deformity, and feeblemindedness The actual hormone. discharged by the gland is thyroxine which appears to act as a stimulant of all metabohe processes, its effect is slowly developed and long lasting, and it is not in general possible to discern it by applying thyroxine to isolated tissues or organs Thyroxine is too insoluble to be given advantageously by mouth, but it is not destroyed during digestion, so that preparations of desiccated thyroid gland, given orally, are cheap aud fully active Methods of biological assay depend chiefly on meas arements of the metabolism of myvedema patients, or of small animals, less accurate methods depend on the fact that thyroid substance accelerates the



Fig 211 EFFECT OF THYROXINE ON GROWTH OF A CRETIN

The illustration shows the same child in the same dress before and after receiving thyroxine for one year Increase in height 6 inches

From kendall Ind Eng Chem 17 520 (19°a)

that thyroid substance accelerates the metamorphosis of tadpoles Thyroxine may also be determined chemically by its iodine content, but it must be remembered that thyro-globulin also contains the more soluble, physiologically inactive amuo acid,

Experiments with the radioactive isotopes of iodine indicate that the gland, unless previously saturated, takes up iodide from the blood for the synthesis of dioadoty rosine and thence of thyroxine. These reactions can be inhibited by treatment with large doses of sulfa drugs or with thoures.

thiour ieil, and related compounds, which thus gradually product signs of hypothyroidism accompanied by gotter. Huouracil is being used experimentally in the treatment of Grave's disease, the administration of iodine also produces a remission, which though often temporary is valuable in preparation for operation, and is due to retention of colloid within the gland.

The uptake of radioactive include (in the form of NaI or KI) can be followed by placing "counters" on the skin above the thyroid, and this has been used as a test of thyroid activity. In larger doses radioactive nodides have been used to control thyroid careinoma and its metastases, since the intensity of radioactivity becomes much higher in the thyroid cells, normally control to the control thyroid cells, normally control to the control thyroid cells, normally control to the contr

mal or cancerous, than elsewhere in the body

THE PARATHYROIDS

The parathyroid glands are small, compact bodies, usually four in num ber and closely apposed to or within the thyroid. If they are removed, the level of calcium in the blood serum falls, this in turn eauses neuromuscular hyperexcitability, tremors of the skeletal muscle, and panting which produces alkalosis With calcium deficiency, this leads to violent tonic and clonic convulsions in which the animal sooner or later dies from arrest of respiration, symptoms best seen in the dog Acid extraction of ox parathyroids yields extracts which maintain parathyroidectomized dogs alive and free from tetany, and which in normal dogs produce an elevation of serum calcium The (Hanson) unit is 0 01 × the amount required to raise the serum calcium of normal, 20 kg dogs by 1 mg per 100 ml, though larger mercases can be more accurately measured A large single injection may accelerate the renal excretion of phosphates and raise the serum calcium to twice its normal level in 12 to 15 liours, usually without disturbance to the health of the animal but if the calcium is maintained at a high level hy repeated injections the blood becomes concentrated and the circulation sluggish, kidney function fails and hemorrhage occurs in the gastrointestinal tract The calcium mobilized by the parathyroid hormone comes from the skeleton, where it is probable that the hormone stimulates the formation and activity of the ostcoelasts After prolonged treatment or in the presence of a parathyroid tumor, the boncs are demineralized and become fibrous and eystie (hyperparathyroidism osteitis fibrosa) There is some tendency for the body to lose its responsiveness to parathyroid extracts this is particularly well seen in the rat The extracts being obtained by rather drastic procedures, there is no certainty that the principle they contain is the unaltered natural hormone, it is of protein nature17 and destroyed by digestive enzymes

EXPERIMENTS ON THE PARATHYROIDS

1 Preparation of Parathyroid Extract Method of Collip Place the fresh ox parathyroids, from which the visible fai and connective tlasue have been removed, in a small flask or test tube cover with an equal volume of 5 per cent hydrochloric acid, and keep in a boiling water bath for 1 hour Allow to cool, and remove the congesied fat mechanically Then make the solu

¹⁷ Ross and Wood J Biol Chem. 146 49 (1343)

tion faintly alkaline with sodium hydroxide and add acid to the point of maximum precipitation of protein, which is just barely acid to litmus. Filter and dissolve the precipitate in weak alkali, and reprecipitate as before. Filter again and combine the filtrates, adding 0.1 per cent tricresol as preservative and keep the combined filtrates, which constitute the desired extract. In an icebox until ready to use.

2. Effect of Parathyraid Extract. Draw 8 to 10 ml. of blood from the saphenous or other suitable vein of a dog weighing at least 10 kg. Save this for determination of serum calcium (p. 644). Inject subcutaneously, every 3 hours for the next 9 to 12 hours, 10 Hanson units of commercial parathyroid extract (or a quantity of extract equivalent to 1 ox parathyroid) per kg. of body weight. Draw blood samples after the last injection, and again 12 hours later. Note the occurrence of depression and anorexia, the increased concentration of the blood, and the rise in serum calcium.

THE ANTERIOR PITUITARY OR HYPOPHYSIS

The anatomy of the pituitary gland, or hypophysis ccrebri (Fig 212) is complex: the pars nervosa is a downgrowth from the hypothalamic region

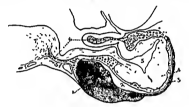


FIG 212 HYPOPHYSIS OF CAT.

1, Pars anterior; 2, hypophysial cleft; 3, pars intermedia; 4, pars nervosa; 5, infundibular cavity; 6, pars tuberalis, 7, optic chiasin.

From Best and Taylor Physiological Basis of Medical Practice. Courtesy, William Wood & Co

of the hrain, and consists of modified nervous and glial cells, while the rest of the organ is derived from Rathke's pouch, an upgrowth from the roof of the mouth. This gives rise to the pars tuberalis, which surrounds the stalk of the pars nervosa like a collar and spreads along the base of the brain, to the pars intermedia, closely apposed to the anterior surface of the nervosa and usually separated by a cleft (the cavity of the embryonic pouch) from the large, glandular pars anterior which lies in front and on either side of it. The gland can be broken in two at the cleft, leaving the untermedia and tuberalis adherent to the nervosa as a complex called the posterior lohe. The whole organ rests in a more or less fitting cavity in the sphenoid hone, the sella turcae.

Removal of the whole organ or of the anterior lobe alone produces the following effects (best seen in the rat): cossation of growth; atrophy of the ovaries or testes, which cease to function in producing germ cells and endocrine organs, atrophy of the thyroid, the reduction of thyroid activity lowering the metabolic rate, atrophy of the adrenal cortex, which may be the cause of an increased susceptibility to toxins, infections, etc., and of a tendency toward dissipation of earhohydrate stores and toward bypoglycemia, and cessation of lactation, if in progress. The administration of suitable extracts, or implantation of anterior-lobe tissue, more or less corrects these deficiencies It is uncertain how many distinct active principles are present in such extracts, but a number are widely recognized and have been separated in pure or nearly pure state. All are of protein or polypeptide nature, are destroyed by digestive enzymes, and are relatively unstable to heat There may be interspecific differences in chemical composition rather than in physiological properties, and this may explain the observation that animals treated with anterior-lobe extracts over long periods tend to become unresponsive to them Clinically, overactivity of the anterior lobe is associated with giantism and aeromegaly (often accompanied by symptoms of diabetes and of hyperthyroidism) and congenital underactivity with dwarfism and infantilism. In Simmonds' disease the gland is almost wholly destroyed

The hormones generally recognized as separate entities are (1) the growth (somatotropic) hormone, capable of producing accelerated growth in young animals, especially growth in length, and active in metabolism in a "contra-insulin" sense, so as to cause severe hyperglycemia in susceptible animals and sufficient damage to the islets of Langerhans to produce a permanent "metahypophyseal" diabetic state, (2) the thyrotropic hormone, stimulating the thyroid gland to activity (reflected in increased beight of the epithchial cells, discharge of iodine-containing thyroxine, and bence raised metabolic rate) and producing exophthalmos, (3) the adrenocorticotropic hormone (ACTH), stimulating the secretion of glucocorticoids (see p 758), (4) prolactin, the "lactogenic" bormone, which stimulates milk secretion in fully developed mammary glands, and the glandular development of the crop seen in pigeons of both sexes while rearing their young, and which also has the luteotropic action of main taining the structure and endocrine function of the corpora lutea, finally, there are the gonadotropic hormones, differentiated as (5) "follicle-stimu lating" (FSH) and (6) "luternizing" or "interstitial cell-stimulating (LH or ICSH), which together incite the sex glands to both germinal and endocrine activity (for example, they produce, when injected into imms ture female rats and miee, premature ripening of follicles, ovulation, formation of corpora lutea, and discharge of ovarian hormones which secondarily affect the uterus and vagina) A substance similar to the luternizing hormone is produced by the human placenta during pregnancy and is termed chorionic gonadotropin (CG) or prolan, it is excreted in the urine, and the Aschheim-Zondek, Friedman, and other similar tests, based on this fact, are the most accurate means now available for diagnosing pregnancy in its early stages These tests also serve for the diagnosis of chorocepithelioma and hydatidiform mole, in which extremely large amounts of gonadotropin are exercised The serum of pregnant mares (PMS) contains yet another gonadotropin, which though of chorionic origin has a closer resemblance to FSH, it is not excreted in the urine

Gonadotropic substances (FSH and LH) of pituitary origin may appear in the urine when the sex glands are removed or have ceased to function; so, too, removal or inactivation (e.g., hy thiouracil) of the thyroid leads to overproduction of thyrotropic hormone. It is evident that the secretory activity of the anterior lohe is to some extent regulated hy the concentration in the hlood of the hormones (estrogens, progesterone, androgens, corticoids, thyroxine) produced hy its "target organs," so that a self-regulating or "servo" mechanism is set up. These hormones probably influence a center in the brain which controls the anterior lobe by a mechanism partly neural and partly humoral, and which cau also be influenced by incoming nervous stimuli (e.g., those arising in the act of mating or the act of suckling) and hy higher cerebral centers.

THE POSTERIOR PITUITARY

The posterior lobe contains principles which may he extracted from the acctone-dried gland with dilute acetic acid; Pitutirin is a commercial extract of this type. It contains assopressin (Pitressin) which, on the one hand, affects the circulation, causing constriction of the capillaries leading to extreme pallor and a tendency to raise blood pressure—offset, however, by an unfavorable action on the heart (coronary constriction)—and, on the other hand, checks the secretion of urine, especially by postpoming or abolishing the diuresis normally induced by administration of fluids. Oxylocin (Pitoicn) is also present in Pitutini but may be almost completely separated from Pitressin; it causes contraction of the smooth muscle of the uterus (except in the presence of an active corpus luteum), and is used therapeutically in the late stages of labor. The extracts also

affect the intestinal and bronchial musculature, and cause squeezing-out of milk from the ducts of the mammary gland

The total synthesis of an octapeptide amide having the hormonal activity of ovytocin by du Vigneaud and associates¹⁸ has established a landmark since it is the first polypeptide hormone to bave heen produced in the lahoratory. Its molecular weight is approximately 1000, the molecule consists of eight amino acid equivalents (leucine, isoleucine, tyrosine, proline, glutamic acid, aspartic acid, glycine, and cystine) and three equivalents of ammonia, the structure being in part cyclic. (See p. 775)

The synthesis of the vasopressor antidiuretic hormone of the posterior

pituitary hy the Cornell investigators appears imminent 19

If the posterior lohe is extripated, or if the nerve fihers which run into it from the nucleus supraopticus are severed, there is a permanent increase in urine flow and in water consumption (diahetes inspiritus). The posterior lohe principles arise in the pars nervosa and are polypeptides, they are unstable in alkaline solution and inactive by mouth. There is evidence that, as ordinarily obtained, they may be hreakdown products of a much larger molecule possessing both pressor and oxytocic activity. The pars intermedia, which is embry ologically part of the anterior lohe, produces a substance (intermedia) which causes expansion of pigment masses in certain cells (melanophores, crythrophores) in the skin of lower vertebrates. A frog from which the pituitary is removed is rendered perma whether this substance has any function in mammals.

EXPERIMENTS ON THE PITUITARY

I Preparation of Anterior Lobe Extract Method of Evans ** Bowine anterior lobes, as fresh as possible, are ground in a mortar with twice their weight of sand, 2 ml of water are added for every g of gland. The volume of the mixture is measured and three eighths of this quantity of 0.2 N Noil is added, the material is kept for 12 hours in the icebox, the supernatiant fluids decanted off and neutralized to phenol red with 0.2 N acetic acid Crude extracts of this type contain maximal amounts of the several hormones extracts of this type contain maximal amounts of the several hormones extracts of this type contain maximal amounts of the several hormones anterior lobes in the lyophilizing apparatus, when dry the tissue crumbles to a light, relatively stable powder, from which uniform suspensions are easily prepared it may be advantageous to add small amounts of penicillin and streptomycin, so as to lessen the risk of abscesses forming at the size of light.

is allowed to stand overnight. The precipitated henzoic acid, upon which the active material is adsorbed, is filtered off with suction and dissolved in a volume of acetone equal to the amount originally added. A small flocculent precipitate of acetone-insoluble material contains the active principle and is separated by decantation and centrifuging and thoroughly washed with acetone. Small additional quantities may be obtained by repeating the henzoic acid adsorption on the urine filtrate. The active principle is extracted from the combined precipitates by three treatments with distilled water, centrifuging after each extraction, and using not more than 25 mi. of water per liter of urine. The water-insoluble residue is discarded. Other useful methods involve adsorption upon Lloyd's reagent²³ or precipitation with tungstic acid.²¹ All these methods may be used to concentrate the activity in urines other than those of pregnancy, when the gonadotropic substance is too dilute to be determined by direct biological assay of the urine.

- 3. Detection of Chorionic Gonodotropin (Prolon): Pregnoncy Test of Aschheim and Zondek.24 Five haby female mice, 6 to 8 g. in weight and 3 to 4 weeks oid, are injected with the urine to be tested. This should be taken from the first sample passed in the morning, and toxic substances may be removed from It by shaking with ether. Each mouse is given 6 injections of 0.5 ml, of tilis urine, 3 doses on the first day and 3 on the second, and the animals are killed 96 hours after the beginning of the test. The reaction sought consists of three parts; (i) Formation of large ovarian follicles and precoclous appearance of estrus; (ii) hemorrhagic follicles, easily seen under a lens as deep clear red spots; (III) formation of corpora iutea, visible under a lens, Reaction I by itself is insufficient to establish a diagnosis of pregnancy. The dependability of the reaction is at least 98 per cent. It usually becomes positive a few days after the first missed menstrual period; in the ensuing month, the concentration of the active principle rises to a well-marked maximum. Very high concentrations occur in the presence of tumors of placental tissue (bydatid moie or chorionepitheijoma).
 - 4. Friedman Pregnancy Test.¹¹ This test is more rapidly completed, is equally accurate, and probably more convenient where tests are made only occasionally. It is hased on the fact rhat rabbits ovulate only when their ovaries are specifically stimulated, as they are alter mating. A mature female rabbit, which must have heen isolated in a cage hy Itself for at least three weeks previously, is given 10 ml. of urine to he tested by injection into the marginal ear vein. The ovaries are examined 24 hours later; a positive reaction is marked by reddish protrusions or by recent hemorrhage in the folicies. The abdomen may he opened and the ovaries inspected under acesthesia, so that the rabbit may he used again some weeks later, when the corpora lutea formed have disappeared.

THE KIDNEY

When the kidneys or the arteries supplying them are compressed, there follows a sustained rise in systemic blood pressure, resembling the clinical entity of essential bypertension; the phenomenon is independent of nervous connections and of changes in everetory function. It is ascribed

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to the secretion into the blood of a globulin renin, which appears to be a proteolytic enzyme and to vary in composition from species to species. The substrate for renin is a fraction (hypertensinogen or "activator") of the serium globulins, and the product of the reaction a heat-stable, dialyzable vasoconstrictor, probably a polypeptide, called hypertensin or angiotomin, this in turn is slowly destroyed by a substance ("inhibitor," angiotomic, or hypertensinase) present in normal blood and tissues

THE THYMUS AND PINEAL

Endocrine function has frequently been ascribed to these organs, but has not been established, extirpation, even in very young animals, bas no specific systemic effects

OTHER HORMONES

The first hormone whose function was clearly established (by Bayliss and Starling in 1903) was secretin. This is a substance which is liberated from the walls of the duodenum when acid ebynee enters the lumen, it is carried by the blood to the pancreas, where it stimulates the secretion of panereatic puice. Tests on the most highly purified secretin show it to be a polypeptide. See also p. 303

Less well established are cholecystolinin, helieved to arise in the same source and to stimulate contraction of the gallbladder, enterogastrone, believed to inhibit the secretory and motor activity of the stomach when fat is present in the intestine, and gastrin, which physiologically resembles instamine, and may evoke secretion of hydrochloric acid by the fundus of the stomach when the pyloric region is stimulated by the presence of partially digested food

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27

Urine: General Characteristics of Normal and Pathological Urine

Secretion of Uririe. The problem of the mechanism of the formation of urine by the kidney has occupied the attention of investigators for many years, and has produced many conflicting theories. The filtrationreabsorption theory of Cushny has been the basis of the modern theory, and the work of Starling, Marshall, Richards and their collaborators and of other investigators has led to a more satisfactory conception of the

processes involved

Marshall has summarized the known facts relating to the formation of urine hy the mammalian kidney in the following working hypothesis "All of the non-colloid constituents of the plasma are eliminated by filtration through the glomerulus, water, chloride, bicarbonate potassium (possibly phosphate, uric acid, and other bodies) are reabsorbed during the passage of the filtrate along the tubules, urea and sulfates are also reabsorbed to some extent under certain conditions but not as actively as chloride ammonia, hippuric acid, and possibly other bodies are formed in the renal cells and secreted, while certain foreign substances when present in the organism (phenol red) and possibly substances occurring in small amounts in the organism are secreted after a preliminary concentration in the tubule cells. The amount of glomerular filtrate on this hypothesis is con sidered sufficient to account for all of the sulfate and urea eliminated but not sufficient to account for ammonia certain dyes (phenol red, etc) and other bodies which are concentrated to a much greater extent than sulfate by the kidney "

By direct eatheterization of the glomerular urine of the frog and com parison of its composition with simultaneously collected bladder urine Wearn and Richards formulated a hypothesis illustrated diagrammati cally in I ig 213 It will be noted that the chief difference from Marshall s conception is in the site of dye sccretion these workers baving found small amounts of dye in the glomerular filtrate although they admit much of the neutral red exerction is earned out by the tubules They also found the polysaccharide mulin to be exercted by the glomerulus, indicating the presence of pores of considerable size in the membrane

Reabsorption of water which may be as high as 97 per cent, appears to occur in both the proximal and distal tubules. Chloride is absorbed prefer entially in the distal tubule, where also takes place the reabsorption of bicarbonate and acidification of the urine, probably through excretion of

hydrogen ions 'Reabsorption of glucose occurs in the proximal tubule, cluefly through esterification with phosphate. The substance phlorizin prevents this esterification and bence permits glucose to pass into the urine ("phlorizin diabetes"). Nitrogenous waste products appear in the urine either through failure of reabsorption from the glomerular filtrate or through actual secretion by the tubular cells or both Ammonia appears definitely to be secreted by the tubular cells, some urea may arise in this manner, and there is evidence for the tubular secretion of creatinine by man and Necturus but not by the frog. Thus species differences must be considered in interpreting kidney action.

Volume. The volume of urme evereted by normal individuals during any definite period fluctuates within very wide limits. The total volume of glomerular filtrate produced by normal adult kidneys may be 75 to 150 liters in 24 hours. However, after tubular reabsorption the average normal.

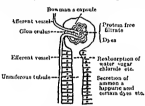


FIG 213 DIAGRAM ILLESTRATING THE FOR

excretion of urine falls within the range of 1000 to 2000 ml. The volume excretion is influenced greatly by the dict, particularly by the ingestion of fluids, and by the ambient temperature, which affects not only fluid intake but also loss of water through perspiration. Strenuous physical exercise causes a diminished output of urine

Certain pathological conditions cause the output of urine for any definite period to depart very decidedly from the normal output. If the output is increased the condition is termed polyura, whereas a dimmished exerction is termed oligina. Among the pathological conditions in which the volume of urine is increased above normal are the following diabetes mellitus, diahetes insipidus, certain diseases of the nervous system, contracted kidney, amyloid degeneration of the kidney, and in convalescence from acute diseases in general. Many drugs such as calomel, digitalis, acetates, and sulcylates also increase the volume of the urine excreted. A decrease from the normal is observed in the following pathological conditions acute nephritis, diseases of the heart and lungs, fevers, diarrhea, and vomiting

Color. Normal urine ordinarily possesses an amher yellow that, the depth of the color being dependent in part upon the density of the fluid

¹ Pitts and Mexander Am. J Physiol 144 239 (1945)

The color of normal urine is due principally to a pigment called urochrome Traces are generally present of urocrythrin, a red pigment possibly derived from the melanins, coproporphyrins, and uroporphyrin, iron-free reddish pigments resulting from the metabolism of heme In porphyria, a disorder of porphyrin metabolism, the porphyrin content is increased In certain other pathological urines a chromogen (indoleacetic acid) may be present. This yields urorosein, a reddish pigment, when the urine is strongly acidified (see p 852) Uroehrome is said to be a compound of urobilin and urobilinogen with a peptide substance The amount excreted per day by an adult man has been estimated at 7 3 mg, being very constant for a given individual Under pathological conditions or after the administration of various drugs or antiseptics, the color of the urine may vary in intensity from an extremely light yellow to a very dark brown or black or may even assume the color of the drugs or their degradation products Vogel has constructed a color chart which is of some value for purposes of comparison. The nature and origin of the chief variations in the urinary color are set forth in tabular form on P 783

Turbid Urine. Normal urine is ordinarily perfectly clear and transparent when voided Oo standing for a variable time, however, a cloud (nubecula) consisting principally of nucleoprotein or nucoid (see p 810) and epithelial cells forms. A turbidity due to the precipitation of phosphates is normally noted in urine passed after a hearty meal. The urine obtained two to three hours after a meal or later is ordinarily free from turbidity. Permanently turbid urines ordinarily arise from pathological cooditions.

Odor. The odor of normal urne is of a faint aromatic type The substances to which this odor is due are not well known but it is claimed by some investigators to be due, at least in part, to the presence of minute amounts of certain volatide organic acids. When the urne undergoes decomposition, e.g., in alkaline fermentation, a very unpleasant ammonical odor is evolved. All urnes are subject to such decomposition if allowed to stand for a sufficiently long time. Under normal conditions the urne very often possesses a peculiar odor due to the ingestion of some certain dray or vegetable. For instance, cubebs copaiba myrtol, saffron, tolu, and turpentine each impart a somewhat specific odor to the urne. After the ingestion of asparagus, the urne also possesses a typical odor attributed to methyl mercaptan (CH, SH) which may, however, exist in urne only as a precursor which yields the mercaptian on heating in acid solution.

Frequency of Urination. The frequency of urination varies greatly in different individuals but in general is dependent upon the amount of fluid in the bladder. In pathological conditions an inflammatory affection of the urinary tract or any disturbance of the innervation of the bladder will influence the frequency. Affections of the spinal cord which lead to an increased irritability of the bladder or a weakening of the sphineter, or any condition lowering the residual capacity of the bladder, will result in increasing the frequency of urination. It often adds in diagnosis

Reaction. The mixed 24-hour urinary exerction of a normal individual ordinarily possesses an acid reaction to litinus. The actual hydrogen-10¹⁰

Color	Cause of Coloration	Pathological Condition
Nearly colorless	Dilution, or diminution of normal pigments	Nervous conditions hy druria, diabetes insipidus, granular kidney
Dark yellow to brown red	Increase of normal, or oc- currence of pathological, pigments Concentrated urme	Acute febrile diseases
Vilky	Fat globules	Cbyluria
	Pus corpuscles	Purulent diseases of the urinary tract
Orange	Excreted drugs	Santonin chrysophanic acid
Red or reddish	Uroerythrin, uroporphyrin coproporphyrin bemoglo bin and myoglobin	Porphyrm bemorrhages, hemoglobinuria, trauma
	Pigments in food (logwood, madder, bilberries, fuchsin)	
Brown to brown black	Hematin	Small hemorrhages
	Methemoglobm	Methemoglobinuria
	Melanin	Melanotic sarcoma
	Hydroquinol and catechol	Phenol poisoning
Greenish yellow, greenish brown, approaching black	Bile pigments	Jaundice
Dirty green* or blue	A dark blue scum on the surface, with a blue de- posit, due to an excess of indigo forming substances	Cholera, typhus, seen espe- cially when the urine is putrefying
Brown yellow to red brown, becoming blood red upon adding alkalis	Substances contained in senna, rhubarh, and cheli donum which are intro duced into the system	

^{*} This dirty green or blue color also occurs after the use of methylene blue in the or ganism

concentration varies over a wide range (pH 48 to 80), the mean being about pH 6 The reaction of the urine represents an equilibrium among a large number of acid and basic constituents, both organic and inorganic, which it contains Although organic acids and bases play a part in producing the normal reaction, this reaction is probably, in the main, dependent upon the relative amounts of the mono- and dibasic sodium and potassium phosphates present The monobasie sodium phosphate (NaH2POi) is acid in reaction, while the dibasic phosphate (Na2HPO4) is alkaline in reaction. The excretion of acid or alkaline phosphate by the kidneys is one of the factors in the regulation of the neutrality of the blood and of the organism in general The acidity of the urine, as determined by titration, runs in general parallel with the hydrogen-ion concentration and seems to be dependent upon the same factors, and in more acid urines mainly on the phosphate content Van Slyke and Palmer have shown that normal men excrete organic acids equivalent to only about 6 ml of 0 1 N acid per kilo in 24 hours. Strenuous physical exercise produces an increase in hydrogen ion concentration and in acid and ammonia output (For further discussion of acidity, see Chapter 31)

The mean acidity in eardiorenal diseases is high—about pH 5 3 as com pared with pH 6, the normal mean In general the acidity tends to be

increased in the greater number of pathological disorders

The composition of the food is perhaps the most important factor in determining the reaction of the urine (see Chapter 34, "Inorganic Metaholism," for the influence of hase forming and acid-forming foods) The reaction ordinarily varies considerably according to the time of day the urine is passed For instance, for a variable length of time after a meal the urme may he neutral or even alkaline in reaction to litmus, owing to the claim of the gastric juice upon the acidic radicals to further the for mation of hydrochlorie acid for use in carrying out the digestry e secretors function. This hypothesis has been verified experimentally. This change in reaction is known as the all aline tide and is common to perfectly healthy individuals. The urine may also become temporarily alkaline in reaction as the result of ingesting alkaline earbonates or certain salts of tartane and citrie acids which ultimately yield bicarbonate within the organism Ingestion of acid fruits (oranges, lemons, peaches, etc.) causes the formation of alkaline urine This is due to the fact that the ash of such fruits is alkaline and when the fruits are combusted in the body bicarbonate is formed On the other hand, bread, cereals, meats, etc., yield an acid ash and an acid urine Certain acid fruits, like cranberries, plums, and prunes, increase urinary acidity. This is due to the fact that these fruits contain considerable quant ties of quinic acid Instead of being oxidized this is converted into hippuric acid, which renders the urine acid III reaction

Normal urine upon standing for some time becomes alkaline owing to the inception of alkaline or ammomacal fermentation through the agency of microorganisms. This fermentation has no especial diagnostic value except in cases where the urine has undergone this change within the organism and is voided in the decomposed state Ammoniacal fermentation is ordinarily due to cystitis or occurs as the result of infection in the process of catheterization. A microscopical examination of such urine (Fig. 214) shows the presence of ammonium magnesium phosphate

crystals, amorphous phosphates, and not infrequently ammonium urate

Occasionally a urine which possesses a normal acidity when voided will upon standing, instead of undergoing ammoniacal fermentation as above described, become more strongly acid in reaction. Such a phenomenon is termed acid fermentation. Accompanying this increased acidity there is ordinarily a deepening of the tint of the urinary color. Sueb urmes may contain acid urates, uric acid, fungi, and calcium ovalate (Fig. 215). On standing for a sufficiently long time any urine which exhibits acid fermentation will ultimately change in reactiou scopic deposits characteristic of such a urine



DEPOSIT IN AMMONIACAL FERUENTATION. a, Acid ammonium urate, b, ammon-

ium magnesium phosphate, c, bacteria owing to the inception of alkaline fermeutation, and will show the micro-

Specific Gravity. The specific gravity of the urine of normal individuals varies ordinarily between 1.015 and 1 025. This value is subject to

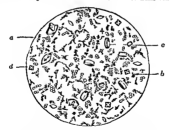


Fig. 215. Deposit in Acid Fermentation. a. Fungus; b. amorphous sodium urate; c. uric acid; d, calcium ovalate.

wide fluctuations under various conditions. For instance, following copious water- or beer-drinking the specific gravity may fall to 1.003 or lower, whereas in cases of excessive perspiration it may rise as high as 1.040 or even higher. Where a very accurate determination of the specific gravity is desired, use is commonly made of the pyenometer or of the Westphal hydrostatic balance. These instruments, however, are not

suited for clinical use. The clinical method of determining the specific gravity is by means of a unnometer (Fig. 216). This affords a very rapid method and at the same time is sufficiently accurate for clinical purposes. The unnometer is always calibrated for use at a specific temperature and

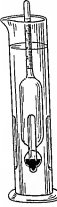


Fig 216 URINOM ETER AND CTL

having a canorated for use at a specific temperature and the observations made at any other temperature must be subjected to a certain correction to obtain the true specific gravity. In making this correction one unit in the third deemal place is added to the observed specific gravity for every three degrees centigrade above the normal temperature and subtracted for every three degrees below the normal temperature. For instance, if m using a uninometer calibrated for 15° C the specific gravity of a urine having a temperature of 21° C is determined as 1018, it is necessary to add to the observed specific gravity 2 × 001 to obtain the real specific gravity of the urine. Therefore the specific gravity at 15° C of a urine having a specific gravity of 1018 at 21° C, would be 1018 + 0002 = 1020.

Pathologically, the specific gravity may be subjected to very wide variations. This is especially true
in diseases of the kidneys. In acute nephritis ordinarily
the urine is concentrated and of a high specific
gravity, whereas in chronic nephritis the reverse con
ditions are more apt to prevail. In fact, under most
conditions, whether physiological or pathological, the
specific gravity of the urine is inversely proportional
to the volume excreted. This is not true of diabetes
mellitis, however, in which the volume of urine is
large and the specific gravity also is high, owing to

the sugar contained in the urine

The total solids normally excreted in the urine may

be roughly calculated by means of Long's coefficient 1c, 26 The solids content of 1000 ml of urine is obtained by multiplying the last two figures of the specific gravity observed at 25° C by 26 To determine the amount of solids excreted in 24 hours if the volume was 1120 ml and the specific gravity was 1018 the calculation would be as follows

- (a) $18 \times 26 = 468$ g of solid matter in 1000 ml of urine
- (b) $\frac{46.8 \times 1120}{1000} = 32.4 \text{ g}$ of solid matter in 1120 ml of unne

Collection and Preservation of the Urine Sample. If any depend able data are desired regarding the quantitative composition of the urine the examination of the mixed excretion for 24 hours is absolutely necessary. In collecting the urine the bladder may be emptied at a given hour, say 8 A v., the urine discarded and all the urine from that hour up to and including that passed the next day at 8 A M, saved, thoroughly mixed, and a sample taken for analysis.

Toluene is a very satisfactory preservative for urine. In using this preservative simply overlay the urine with a thin layer of the toluene Formaldehyde (2 drops per 50 ml of urine) or a bit of camphor or thymol are also satisfactory urine preservatives which do not interfere with the tests for the major urinary constituents

Another satisfactory preservative consists of a 3 2 mixture of hexamethylenetetramine (urotropin) and salicylic acid, which is used in the proportion of 50 mg per 10 ml of urine These substances produce formaldehyde in solution and, it is claimed, do not interfere with any of the usual tests

In certain nathological conditions it is desirable to collect the urine passed during the day senarately from that passed during the night. When this is done, the urine voided between 8 AM and 8 PM may be taken as the day sample and that voided between 8 PM and 8 AM as the night sample

The qualitative testing of urine samples collected at random, except in a few specific instances, is of no particular value so far as giving us any accurate knowledge as to the exact urmary characteristics of the individual is concerned. In the great majority of cases the qualitative as well as the quantitative tests should be made upon the mixed excretion for a 24-hour period as well as upon a night sample as above described

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28

Urine: Physiological Constituents'

Normal unne vanes widely in composition, being influenced by diet and other factors. The following table represents the composition of average normal unne

COMPOSITION OF A TYPICAL NORMAL UBINE

	Daily Excretion in
Constituent	Grams.
Nater	1200 0
oglida	60 0
Crea	30 0
Uric acid	0 7
Hippurie aci l	0.7
Creatiniii	12
Indican	0 01
Oxalic acid	0 02
Mantoin	0 04
\mino acid nitrogen	0 2
Purine bases	0 01
Pi enols	0.2
Chloride as \aCl	12 0
bodium	4 0
Potassium	2 0
Calcium	0 2
Magnesium	0 15
bullur, total, as 5	10
Inorganie sulfates as 5	0.8
Seutral sulfur as 4	0 12
Conjugated a illates as 5	0 03
Pleastain na V	3.37

Urea is the principal end product of the metabolism of protein substances in mammals, amphibia, and clasmobranch fishes. About 80 to 90 per cent of the total nitrogen of human urine is present as urea. The distribution of the mitrogen of the urine among urea and the other nitrogencontaining compounds present depends upon the absolute amount of the total nitrogen excreted A decrease in the total nitrogen excretion is always accompanied by a decrease in the percentage of the total nitrogen excreted as area. By so reducing the protein content of the diet of a normal person as to cause the excretion of total nitrogen to be reduced to 3 to 4 g in 24 hours, only about 60 per cent of this nitrogen appears in the urine as urea Uren is the only one of the nitrogenous excretions which is decreased, relatively as well as absolutely, as a result of decreasing the amount of protem metabolized Folin reported a bospital case in which only 14 7 per cent of the total nitrogen was present as urea and about 40 per cent was present as ammoma Morner had previously reported a case in which but 4.4 per cent of the total nitrogen of the urine was present as urea, and 26.7 per cent was present as ammonia

Urea occurs most abundantly in the urine of man and carnivora and in a manifer amount in the urine of herbivora, the urine of fishes, ampl binans, and certain birds also contains a small amount of the substanc. Urea is also found in nearly all the fluids and in many of the tissues and organs of mammals. The amount excreted under normal conditions by an adult man in 24 hours is about 30 g. The excretion is greatest in amount on a diet of meat, and least in amount on a diet consisting of nonintrogenous foods, this is due to the fact that the urea output is regulated by the protein metabolism. A low-protein diet has a tendency to decrease the metabolism of the tissue proteins and thus the output of urea under these conditions may fall below that observed during starvation. The output of urea is also increased after copious water-or beer-drinking. The increase is probably due primarily to the washing out of the tissues of the urea previously formed, but which had not been removed in the normal processes, and accondarily to a stimulation of protein catabolism.

The formation of urea appears to be a property of the liver exclusively. This has been demonstrated by experiments in which there was complete cessation of urea formation after extripation of the liver Studies with isolated tissues, incubated in thin shees under physiological conditions, have also shown that only liver tissue is capable of synthesizing urea.

The mechanism of urea formation by the liver is still uncertain Early views included formation from free NH₄, presumably liberated from amino acids by oxidative deamination, through the intermediate formation of ammonium carbonate, (NH₀CO₂, or ammonium carbonate, NH₄O CO NH₂ Atteution has shifted to the role of certain amino acids in urea formation because of the studies of Krebs and Henselet These investigators, working with liver slices, showed that the liver could synthesize urea from ammonia and postulated the evistence of a cyclic mechanism including ammonia, carbon diovide, and the amino acids anginine, ornithine, and etiruline The steps in the process, recording to the ornithine cycle theory, are as follows

Thus the primary source of urea according to this theory is the action of the enzyme arginase on the amino acid arginine to produce urea and ornithine. The ornithine acis as a catalyst being converted back to arginine by the reactions shown, taking up aimmonia and carbon dioude.



in the process which ultimately appear as urea. This theory, while attractive is not universally accepted (see Chapter 33). Points in its favor include the well known and powerful arginase activity of liver and the fact that circuling first isolated from watermelon juice is found in small amount in the blood and its synthesis by liver tissue can be demonstrated. Opposed to it is the fact that the experimental demonstration of the orinitime cycle with liver sheets requires the presence of free am

The birret may be dissolved in water and a reddish-violet color obtained by treating the aqueous solution with copper sulfate and potassium hydrovide (see "Birret Test," p. 171). Certain hypochlorites or hypobromites in alkaline solution decompose urea into nitrogen, earbon dioxide, and water. Sodium hypobromite brings about this decomposition as follows:

$$CO(NH_2)_2 + 3NaOBr \rightarrow NaBr + N_2 + CO_2 + 2H_2O$$

According to the Werner hypothesis, the urea is momentarily brominated and the resultant compound hydrolyzed by the alkali and oxidized by hypobromite. Sodium cyanate, sodium nitrate, hydrazine, and CO are formed as by-products, thus accounting for the low nitrogen values obtained in the clinical application of this method.

Soybeans, jack beans, and watermelon seeds contain an enzyme called urease which has the power to decompose urea with the liberation of ammonia. This fact is made use of in the quantitative determination of urea (see Chapter 23). Urease action appears to involve the intermediate formation of ammonium carbamate and not of cyanate.

Urea forms crystalline compounds with certain acids; urea nitrate and urea oxalate are the most important. Urea nitrate, CO(NH₂)₂·HNO₂, crystallizes in colorless rhombic or six-sided tiles (Fig. 218), which are easily soluble in water. Urea oxalate, (CO(NH₂)₂)₂·H₂C₂O₄, crystallizes in the form of rhombic or six-sided prisms or plates (Fig. 219): the oxalate differs from the nitrate in being somewhat less soluble in water.

A decrease in the exerction of urea is observed in many diseases in which the diet is much reduced, in diseases associated with impaired liver function, in some disorders as a result of alterations in metabolism, e.g., myxedema, but most frequently as a result of diminished exerction, as in



FIG 218 UREA NITRATE.

severe and advanced kidney disease. In fact, the determination of the ability of the kidneys to exercte urea (the "urea clearance" test, see Chapter 31) is perhaps the most valuable single clinical index of real function. A pathological increase may result from tissue catabolism in

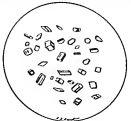


FIG 219 UREA OXALATE.

febrile or wasting conditions. In marked acidosis it may be considerably decreased relative to the total introgen (see * Ammonia " p. 814). Strong solutions of ura have a remarkable solvent effect upon proteins.

such as coagulated protein, and upon starch and other substances

EXPERIMENTS ON UREA

- 1. Isolotion from the Urine, Evaporate 200 ml, of urine in a casserole or evaporating dish over a free flame until the volume is reduced to about 25 ml. Transfer the container to a steam or water hath and continue the evaporation until the residue is semisolid, spreading the material over the sides of the dish to aid in removing as much water as possible. Add 50 ml. of acetone to the residue and stir thoroughly while the acetone is kept cently holling on a previously heated water bath (with the flame turned out) or steam bath. Do not allow more than one-quarter of the acetone to boll off. Because of the inflammable and polsonous nature of acetone vapor, this and further operations should be conducted in the hood and in the absence of a free flame. Filter the hot extract guickly through a small dry filter into a dry 100-ml, beaker. Repeat the extraction of the residue once more with acetone. adding the filtrate to the first portion. Concentrate the combined extracts to a volume of 25 ml, on the steam bath or by placing the beaker in hot water, Chill the concentrated solution with cold water, cover, and allow to stand until crystallization occurs. Filter off the urea crystals, wash them with a little acetone, and allow to dry in air. Examine the crystals under the microscope and compare them with those shown in Fig. 217.
- 2. Melting Point. Determine the melting point of some urea crystals as follows: Into an ordinary melting-point tube, sealed at one end, introduce powdered urea. Fasten the tube to the bulb of a thermometer and suspend the bulb and its attached tube in a small beaker containing concentrated sulfuric acid. Gently raise the temperature of the acid by means of a low flame, stirring the fluid continually, and note the temperature at which the urea begins to melt.
- 3. Crystolline Form. Dissolve a crystal of pure urea in a few drops of 95 per cent alcohol and place I to 2 drops of the alcoholic solution on a microscopical silde. Allow the alcohol to evaporate spontaneously, examine the crystals under the microscope, and compare them with those reproduced in Fig. 217. Recrystallize a little urea from water in the same way and compare the crystals with those obtained from the alcoholic solution.
- 4. Formation of Bluret. Place a small amount of urea in a dry test tube and heat carefully in a low diame. The urea melts at 132° C. and liberates ammonia. Continue heating until the fused mass hegins to solidify. Cool the tube, dissolve the residue in dilute sodium hydroxide solution, and add very dilute copper sulfate solution (see p. 171). The purplish-violet color is due to the presence of hiuret which has heen formed from the urea through the application of heat as indicated. The chemistry of this reaction is shown on p. 791.
- 5. Urea Nitrate. Prepare a concentrated solution of urea by dissolving a little of the substance in a few drops of water. Place a drop of this solution on a microscopical silide, add a drop of concentrated nitric acid, and examine under the microscope. Compare the crystals with those reproduced in Fig. 218.
- 6. Urea Oxalate. To a drop of a concentrated solution of urea, prepared as described in hxp. 5, add a drop of a saturated solution of oxalic acid. Examine under the microscope and compare the crystals with those shown in Fig. 219.

7. Decomposition by Sodium Hypobromite Into a mixture of 3 ml of concentrated sodium hydroxide solution and 2 ml of bromine water in a test tube introduce a crystal of urea or a small amount of concentrated solution of urea Through the influence of the sodium hypobromite, NaOBr, the urea is decomposed and carbon dioxide and nitrogen are liberated The carbon dioxide is absorbed by the excess of sodium hydroxide, while the mitrogen is evolved and causes the marked effervescence observed. This property forms the basis for one of the methods in common use for the quantitative determination of urea Write the equation showing the decomposition of urea by sodium hypobromitee.

It is claimed that all ammon um compounds and all compounds containing the amino [-NH₂] group yield nitrogen when treated with hypobromite as in this test

8 Decomposition by Urease To 5 ml of urea solution in a test tube add i ml of urease solution or a little soyhean or jack bean powder Allow the tube to stand for 10 minutes, heat the contents to boiling, holding moist red and blue litmus papers at the mouth of the tube What do you observe? Note the odor Explain

URIC ACID

Unc acid is found in the urine normally to the extent of about 0 o to 10 g per 24 hours but this amount is subject to wide variations par ticularly under certain detary and pathological conditions. On a purine-free diet the uric acid output may be 0 1-0 5 g per day whereas a high purine diet may yield a daily output of 2 g. Uric acid acts as a weak dibasic acid and forms two classes of salts neutral and acid. The neutral potassium and lithium urates are the most easily soluble of the alkalisalis, the ammonium urate is difficultly soluble. The acid salts are more insoluble and form the major portion of the sediment which separates upon cooling the concentrated urine. The alkaline earth urates are very insoluble. Ordinarily uric acid occurs in the urine in the form of urates and upon acidifying the hquid the uric acid is liberated and deposits in crystalline form.

Ure acid is closely related to the purine bases as may be seen from a comparison of its structural formula with those of the purine bases given on p 202 'ecording to the purine momentature, it is designated 2.6 5 oxypurine Ure acid forms the principal end product of the nitrogenous ctabolism of birds and scal) replies in the human organism it occupies position quantitatively inferior to urea ammonia and creatinine

In man time acid probably results principally from the destruction of actian or other purine material injected as food or from the dismits, rat ig cellular matter of the organism. The une acid resulting from the dist



URIC ACID CRESTALS \ORMAL COLORS
(From Purily after Peyer)

was formerly said to be of exogenous origin, whereas the product of cellular catabolism was said to be of endogenous origin. It is now known that metabolic activities cannot be properly explained on such a simple basis. However, the terms endogenous and exogenous are nevertheless frequently employed. Folin demonstrated that, following a pronounced decrease in the amount of protein metabolized, the absolute quantity of uncaid is decreased, but that this decrease is relatively smaller than the decrease in the total introgen excretion and that the percentage of the uncased introgen, in terms of the total introgen, is therefore decidedly interested.

The enzymatic conversion of the purines adenine and guanine to the intermediates hypoxanthine and xanthine, and of the latter to uric acid by means of xanthine oxidase, is discussed in Chapter 7, "Nucleic Acids and Nucleoproteins" With the exception of man, the higher ages, and the Dalmatian dog, mammaha carry the conversion one sten farther, i.e., to allantoin, through the action of the enzyme uricase Despite the absence of a typical uricase from human tissues, there is no question but that the human organism has considerable ability to destroy une acid, by mechanisms not known Thus uric acid excretion represents a balance between the rate of production and the rate of destruction From experiments on dogs. Mann and associates drew the conclusion that the destruction of unc acid depends on the presence of the liver The extirpation of the liver causes an accumulation of uric acid in the blood and tissues or an increased elimination in the urine if renal activity is maintained. These experiments also indicate that the liver is of considerable importance in the general metabolism of the purines

Using the isotope technique it has been shown that orally administered unce acid is "extensively degraded" to urea, whereas intravenously administered unceacid is exercted essentially unchanged. This latter finding is in agreement with the reported absence of uncase in human tissues.

In hirds the formation of uric acid is analogous to the formation of urea in man. In these organisms it is derived principally from the protein material of the tissues and the food and is formed through a process of synthesis which occurs for the most part in the liver, a comparatively small fraction of the total uric acid excretion of birds may result from nuclear material.

When pure, unc acid may be obtained as a white, odorless, and tasteless powder, which is composed principally of small, transparent, crystalline rhombic plates. Uric acid as it separates from the urine is invariably pigmented, and crystallizes in a large variety of characteristic forms, e.g., dumbbells, wedges, rhombic prisms, irregular rectangular or hexagonal plates, whetstones prismatic rosettes, etc. Uric acid is insoluble in alcohol and ether, soluble with difficulty in boiling water (1 1800) and practically insoluble in cold water (1 39,480, at 18° C). It is soluble in alkalies, alkalicarbonates, boiling glycerol, concentrated stiffuric acid, and in certain organic bases such as ethylamine and piperidine. It is claimed that the uric acid is held in solution in the urine by the urea and disodium hydro-

^{*}Geren Bendich Bodansky and Brown J Biol Chem 183 21 (1950)

gen phosphate present. Une acid possesses the power of reducing cupric hydroxide in alkaline solution and may thus lead to an erroneous conclusion in testing for sugar in the urine by means of Fehling's or Trommer's test. A white precipitate of cuprous urate is formed if only a small amount of cupric hydroxide is present, but if enough of the copper salt is present the characteristic rid or brownish red precipitate of cuprous oxide is obtained. Uric acid does not possess the power of reducing bismuth in alkaline solution and therefore does not interfere in testing for sugar in the urine by means of Boettger's or Nylander's tests.

In addition to being a constant urinary constituent uric acid is present in small amounts in normal human blood as well as in the blood of hirds It is also normally present in the brain, heart, liver, lungs, pancreas, and soleen

Pathologically, the excretion of une acid is subject to wide variation, but the experimental findings are rather contradictory. It may be stated with certainty, however, that in leukemia, hecause of the destruction of nuclear material, the une acid output is increased absolutely as well as relatively to the urea output, under these conditions the ratio between the une acid and urea may be as low as 1.9, whereas the normal ratio, as we have even; is 1.30 or higher An actual output of 12 g of une acid per day has been reported in leukema.

In gout the kidney is said to lose the power of properly eliminating une acid and it collects in the blood in abnormally high concentration. This is accompanied by the deposition of une acid in the joint cartilages and especially in the joint of the big too. Tests have been reported, which "minimize the role of dietary purine in the production of overt gout."

The une acid content of the urine is of importance in relation to the formation of une acid calcult. The administration of alkali carbonates and citrates, or the feeding of base-forming foods by decreasing the acidity of the urine, increases its solvent power for une acid and decreases the hability of formation of this type of calculus.

remains which turns purplish-red after the dish has been cooled and a drop of very dilute ammonium hydroxide has been added. The color is due to the formation of murexide, if potassium hydroxide is used instead of ammonium hydroxide, a purplish-violet color due to the production of the potassium salt is obtained. The color disappears upon warming; with certain related compounds (purine bases) the color persists under these conditions. This is a valuable test for the detection of uric acid calculi.

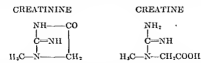
In this reaction the unc acid is oxidized to dialuric acid and alloxan. These two substances condense to form alloxantin This alloxantin reacts with ammonium hydroxide to form purpure acid. The purple color is due to the formation of ammonium purpurate or murexide.



FIG. 220, PURE URIC ACID.

4. Phosphotungstic Acid Reaction (Folin). To 20 ml. of saturated sodium carbonate solution in a small beaker add a small amount of uric acid. Stir the solution until the uric acid has dissolved, then add 1 mi. of Folin's uric acid reagent (see p. 560). A blue color results.

5. Silver Reduction Test (Schiff). Dissolve a small amount of pure urlc acid in sodium carbonate solution and transfer a drop of the resulting mixture to a strip of filter paper saturated with silver nitrate solution. A yellowishbrown or black coloration due to the formation of reduced silver is producedit is calamed that chiordes interfere with this test.



Creatinine. Creatinine is the anhydride of creatine (methylguanidinoacctic acid) and is a constant constituent of normal human urine. Under normal conditions about 1 to 1.8 g. of creatinine are excreted by an adult man in 24 hours. The exact amount of creatinine excreted under ording circumstances depends in part upon the nature of the diet, since any creatinine in the diet is excreted unchanged in the urine Foods such as meat and fish contain significant amounts of performed creatinine, particularly after cooking Creatinine excretion decreases somewhat in starvation The absolute amount of creatinine climinated in the urine on a creatinine-free diet is practically a constant quantity for a given individual and is independent of quantitative changes in the total amount of nitrogen eliminated under these conditions The "creatinine coefficient," which is the daily excretion of creatinine in mg per kg of body weight,4 is an index of this constancy of creatinine climination Endogenous creatinine is apparently the result of some special process of normal metabolism which takes place at a constant rate and almost certainly involves body creatine, since creatine can be converted in the body to creatinine For example, when animals are given tagged creatine the isotopie ratios of tissue creatine and urine creatinine are similar The conversion of creatinine to creatine in the body does not occur

Very little that is important is known regarding the exerction of creatinine under pathological conditions. The creatinine content of the urine is said to be increased in conditions associated with increased tissue catabolism, such as fover The output of creatinine is decreased in dis-



Fig 221 CREATININE

orders associated with muscular atrophy and muscular weakness The greater part of the data relating to the variation of the creatmine excretion under pathological conditions are not of much value since in nearly every instance the diet was not sufficiently controlled to permit the collection of reliable data

According to Mann and associates the changes in the blood creatinine after removal of the liver depend entirely on the kid neys If the kidneys are active the blood creatinine does not change If the kidneys are removed as well or the animal becomes an

urie, the creatinine increases in the blood at the same rate as when only the kidneys are removed The creatine behaves in a similar manner The liver would therefore appear to take no specific part in the creatininecreatine relationship

Creatinine crystallizes in colorless glistening, monoclinic prisms (Fig. 221) which are soluble in about 12 parts of cold water, they are more soluble in warm water and in warm alcobol It forms salts only with strong mineral acids One of the most important and interesting of the compounds of creatinine is creatinine zinc chloride, (C4H7N2O)2ZnCl1,

^{*} Shaffer designates as the creatmine coefficient* the excretion of creatmine nutrogen in mg. per kg. of body weight

which is formed from an alcoholic solution of creatinine upon treatment with zinc chloride in acid solution. Creatinine bas the power of reducing cupric hydroxide iu alkaline solution and of forming an iusoluble cuprous-creatinine salt, and in this way may interfere with the determination of sugar in the urine. Creatinine does not reduce alkaline bismuth solutions and therefore does not interfere with Nylander's and Boettger's tests.

Creatine. Creatine occurs in very small amounts in the urine of normal adults but is found in larger amounts in that of children and of pregnant women. The amount is increased in fasting, in pregnancy, and after high water ingestion. Increased amounts have been found associated with malnutrition and disintegration of muscular tissue, in fever, and in carcinoma of the liver. The presence of readily detectable amounts of creatine in the urine is called creatinuria.

Creatine in the diet does not normally appear as either extra creatinine or creatine in the urine, being apparently incorporated by the body in its total supply of creatine, where it undergoes metabolism at the same rate as that already present. The fact that ingestion of a gram or so of creatine does not normally lead to any change in urinary creatunine or creatine is used clinically in the so-called creature tolerance test, in certain diseases involving muscular tissue chiefly, ingestion of creatine is followed by the appearance of extra creatine in the urine, i.e., the patient's tolerance for creatine is decreased. Creatine occurs in all tissues, but in much greater amount in the muscular tissues than elsewhere. The average adult human hody contains about 100 g. of creatine, largely if not entirely in the form of phosphocreatine.

Creatine is synthesized in the animal hody. The mechanism of this synthesis resisted elucidation for many years, finally yielding to the application of isotopes and the tissue-slice technique, through the work of Schoenheimer, du Vigneaud, Borsook, and their collaborators. While other reactions may not be entirely excluded, these investigators have shown that the major processes in the biological synthesis of creatine are as follows:

I hus the synthesis involves the amino acids glycine and arginine in the reaction of transamidination and a source of labile methyl groups in the transmethylation reaction the latter may involve either the amino acid methionine or the substance choline. In this connection it is of interest that while glycine is readily synthesized by man it cannot be synthesized by the chick and is ordinarily required in the diet but may be replaced there by creatine Of interest also is the clinical observation that in myasthenia gravis the feeding of glyeme leads to an increased excretion of creatine The intermediate compound guanidinoacetic acid is found in normal urine to the extent of about 20 mg per day

Creatine is fairly soluble in water but the aqueous solution is unstable the creature being gradually transformed to its anhydride creatinine by ring closure after the splitting off of water This process is accelerated by heat and acid and this is the basis of the usual procedures for the detection or determination of ereatine the ereatine being converted into creatinine and tested for as the latter compound. There is no satisfactor) direct test reaction for creatine itself

EXPERIMENTS ON CREATININE AND CREATINE

1 Preparation of Pure Creatinine from Urine (Folin Benedict) To 10 liters of undecomposed urine in a large precipitating jar add with stirring a hot solution of 180 g of pictic acid in 450 ml of boiling alcohol Allow to stand overnight and syphon off the supernatant fluid Pour the residue upon a large Buchner funnel drain with suction wash once or twice with cold saturated picric acid and suck dry Treat the dry or nearly dry picrate in a large mortar or evaporating dish with enough concentrated liGl to form a moderately thin paste (about 60 ml of acid for each 100 g of picrate) and stir the mixture thoroughly with the pestie for 3 to 5 minutes Filter with suction on a hardened paper and wash the residue twice with enough water to cover ir sucking as nearly dry as possible each time Transfer the filtrate to a large flash and neutralize with an excess of solid magnesium oxide (the heavy variety is best) Add this oxide in small portions with cooling of the flask under running water between the additions Neutralization of the acid will be indicated by a bright jellow color of the mixture or litmus paper may be used to test it Filter with suction Wash the residue twice with water immediately add a few ml of glacial acetic acid to the filtrate to make it strongly acid Neglecting any precipitate that may form dilute the solution with about 4 volumes of 95 per cent alcohol After 15 minutes filter off the light precipitate which forms Treat the final filtrate with 30 to 40 ml of o per cent zinc chioride Stir and let stand overnight in a cool place Pour off the supernatant liquid and collect the creatinine zinc chloride on Buchner funnel wash once with water then thoroughly with 50 per cent alcohol finally with 95 per cent alcohol and dry A nearly white light cry alal line powder should be obtained The yield should he 90 to 95 per cent of the original creatinine (usually about 1 5 to 1 8 g of creatinine zinc chloride per liter of urine

Recrystallize the creatinine zinc chloride by treating 10 g with 100 ml of water and about 60 mi of normal sulfuric acid heating the mixture until a cless solution is obtained Add about 4 g of purified animal charcoal continue boiling for about a minute filter with suction through a small Buchner funnel pouring the filtrate back on the filter 3 or 4 times until it runs through perfectly colorless Wash the residue with hot water and transfer the total filtrate to a heaker and while hot treat with a little strong zinc chloride solution (3 ml.) and with ahout 7 g. of potassium acetate dissisolved in a little water. After 10 minutes dilute with an equal volume of, alcohol, and allow to stand in a cold place for some hours. Filter off the crystalline product and examine under a microscope (see Fig. 222). To remove, the small amount of potassium sulfate which it contains stir up with its

weight of water, filter, wash with a little water and then with alcohol. The preparation should be snow white.

Yield, 85 to 90 per cent.

For the decomposition of the creation zinc chloride Gaehler suggests the following modification of the original procedure, Place 32 g, creatinine zinc chloride in a pressure bottle (a citrate of magnesia hottle will serve) and add 225 to 250 ml. of concentrated ammonia. Close the hottle and heat in a water bath at 70 to 80° C., sbaking to effect solution. Cool quite rapidly to room temperature and then in a saltice bath. Pure creatinine crystallizes out, Filter. Wash with Ice-cold ammonia, then with acctone, and dry. The yield should be 75 per cent (15 g.

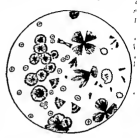


Fig. 222. Creatinine Zinc Chloride (Salkowsei).

of creatinine). The product is perfectly pure and can he used as a standard in the quantitative determination of creatine and creatinine. See the chapters on quantitative analysis of urine and blood.

- Nitroprusside Test (Weyl). Take 5 ml. of urine ln a test tube, add a few drops of sodium nitroprusside, and render the solution alkaline with NaOH solution. A ruby-red color results which soon turns yellow. See Legal's test for acetone. Chapter 29.
- 3. Nitroprusside-Acetic Acid Test (Salkowski). To the yellow solution obtained in Weyl's test above, add an excess of acetic acid and apply heat. A green color results and is in turn displaced by a blue color. A precipitate of prussian blue may form.
- 4. Picric Acid Reaction (Jaffé). Place 5 mi. of urine in a test tube, add an aqueous solution of picric acid, and render the mixture aikaline with NaOH solution. A red color is produced which turns yellow if the solution be acidified. Glucose gives a similar red color hut only upon the application of heat. This color reaction observed when creatinine in aikaline solution is treated with picric acid is the hasic principle of Folin's colorimetric method for the quantitative determination of creatinine (see Chapter 3) and is due to the formation of a red tautomer of creatinine picrate. The production of this tautomer is "dependent upon the formation of a salt, a keto-enoi change within the creatinine molecule, and a change in the picric acid molecule involving the hydrogens in the meta positions and, probably, all three nitro groups."
- Preparation of Creatine. Creatine may be prepared from creatinine zinc chioride by decomposition with calcium hydrate, the process being one of hydrolysis (Benedict).

100 g of creatinine zinc chloride are treated with about 700 mi of water in a large casserole and the mixture heated to boiling, 150 g of pure powdered calcium hydrate are then added, with stirring, and the mixture boiled gently for 20 minutes (with occasional stirring) The hot mixture is then filtered with suction, the residue being washed with hot water The filtrate is then treated with hydrogen sulfide gas for a few minutes and poured through 2 folded filter to remove the zinc. The filtrate is acidified by the addition of about 5 mi of glacial acetic acid and bolied down rapidly to a volume of about 200 ml. This solution is allowed to stand overnight, preferably in a cool place The next day the crystallized creatine is filtered off with suction, washed with a very little cold water, and then thoroughly washed with alcohol and dried This product is then recrystallized by dissolving in about 7 times its weight of boiling water and allowing the solution to cool slowly and stand for some hours. This product should be perfectly pure creatine if necessary it can be recrystailized with very little loss. The crystallized product should be filtered off, washed with alcohol and ether, and dried in air for about half an hour Thus obtained, the creatine contains water of crystallization which it loses very readity upon exposure to air To prepare creatine which can be weighed with absolute exactness it is necessary to dehydrate this product by heating for some hours at about 95° C

The yield in this process is about 18 g of recry stallized creatine, and about 55 g of creatinine zinc chiloride recovered. Longer boiling with time does not bring about a greater yield, for after the 20 minute point creatine is decomposed aimost exactly as fast as it is formed.

Examine the crystals of creatine under the microscope and compare with liliustration in Chapter 10, "Muscular Tissue, in which may be found other creatine tests

ETHEREAL SULFATES

The most common ethereal sulfates found in the urine are phend sulfure acid, p-cresolsulfure acid indox)sulfure acid and skatoxys sulfure acid Pyrocatecholsulfure acid also securs in traces in human urine. The total output of ethereal sulfure acid (as S) varies ordinarily from 0.04 to 0.1 g for 2.1 hours and comprises o to 10 per cent of the total sulfur. In health the ratio of ethereal plus neutral sulfate to morganic sulfate is about 1.10. These ethereal plus neutral sulfate to morganic from the phenol cresol indole and skatole formed in the putrefaction of protein material in the intestine. The phenol passes to the liver where part of it is conjugated to form phenol potassium sulfate and appears in this form in the urine whereas the indole and skatole undergo a preliminary oxidation to form indoxyl and skatoxyl respectively before their conjugation and climination. (See Chapter 20)

It was formerly generally considered that each of the ethereal sulfure acids was formed principally in the putrefaction of protein material in the intestine and that therefore a determination of the total ethereal sulfure acid content of the urine was an index of the extent to which these putrefactive processes were proceeding within the organism Folin however, showed that the ethereal sulfure acid content of the urine did not afford an index of the extent of intestinal putrefaction, since these compounds arise only in part from putrefactive processes. He claimed

that the ethereal sulfurie acid everetion represented a form of sulfur metabolism which is more in evidence when the diet contained little or no protein. The ethereal sulfurie acid content of the urine diminishes as the total sulfur content diminishes, but the percentage decrease is much less. Therefore, relative to the total sulfurie acid content, the ethereal sulfuric acid content is inecessed, although the total sulfuric acid content is diminished. The indoxylsulfurie acid (indican) content of the urine does not originate to any degree from the tissue catabolism of protein material hut arises in great part from the action of intestinal putrefactive organisms on tryptophan (see Chapter 20). The excretion of indoxylsulfurie acid

which occurs in the urine as the potassium salt

may alone he taken as a rough index of the extent of putrefactive processes within the intestine and is clinically the most important of the ethereal sulfurne acids. Under normal conditions, from 10 to 20 mg, of indican are excreted per day. The variations are due mainly to diet, a high meat diet causing an increase and a carhohydrate diet a decrease. Pathologically, the greatest increases are found in disorders involving increased putrefaction and stagnation of intestinal contents. Bacterial decomposition of body protein as in gangrene, point pus formation, etc., gives rise to an increased indican excretion.

Phenols are excreted chiefly in the conjugated form. The phenol output tends to vary directly hut not proportionally with the protein ingestion. The total phenol excretion of normal men on an ordinary mixed diet averages around 0.2 g. per day.

TESTS FOR INDICANS

1. Jafjé's Test. Nearly fill a test tube with a mixture composed of equal volumes of concentrated HCl and the urine under examination. Add 2 to 3 ml. of cbloroform and a few drops of a calcium hypochlorite solution, place the thumb over the end of the test tube and rock the tube back and forth, inverting at least 10 times. The chloroform is more or less colored, according to the amount of indican present. Ordinarily a blue color due to the forma-

⁵ The urms should always be examined fresh if this is possible. In any event formaldehyde should never be used as a preservative for urmes which are to be examined for indican by means of any test involving by pochlorite or potassium permanganate. The formaldehyde through its reducing power lowers the oridinal efficiency of the mixture. The formation of forme and from the aldebyde may also interfere.

tion of indigo blue is produced, less frequently a red color due to indigo red may be noted.

Repeat this test on some of this same urine to which formaidehyde has been added. Is there any variation in the reaction from what you previously obtained?

The following represents the reaction

2. Obermayer's Test. Nearly fill a test tube with a mixture composed o equal volumes of Obermayer's reagent' and the urine under examination Add 2 to 3 ml. of chloroform, place the thumb over the end of the test tube and rock the tube back and forth, inverting at least 10 times. How does thi compare with Jaffe's test?

HIPPURIC ACID

This acid occurs normally in the unne of both carnivora and herbivora, but it is much more abundant in the urine of the latter. It is formed by the union of benzoic acid with glycine according to the following reaction

In the dog this conjugation takes place exclusively in the kidneys, but in man and in the rabbit it is probable that the liver is the main site of this synthesis. Climically the ability of an individual to synthesize hippure and after administration of a test dose of benzoate is used as an index of liver function. Glyenie is readily produced by the body from the hydrolysis of dietary or tissue protein, or may be synthesized from certain other amino acids, e.g., serine Benzoa card originates ordinarily either from intestinal putrefaction, i.e., from the abnormal metabolism of tyrosine and phenylalanine, or from the food. Many vegetables, fruits, and grasses contain small quantities of preformed benzoa card and larger amounts of quine acid and other compounds which in the body are converted to benzoa card. It has been found that approximately 2.g. of benzoa card are exerted as hippuric acid after eating 250 g. of pruines. The average

See Appendix.

excretion of hippuric acid by an adult man for 24 hours under normal conditions is about 0.7 g. Hippuric acid crystallizes in needles or rhombic prisms (see Fig. 223), the particular form depending upon the rapidity of crystallization. Pure hippuric acid melts at 187° C. It is easily soluble in alcohol or hot water. It is sufficiently soluble in ether to allow its extrac-



FIG 223 HIPPURIC ACID

tion from aqueous solution with this solvent (For the quantitative determination of hippuric acid, see Chapter $\bf 31$)

EXPERIMENTS ON HIPPURIC ACID

- 1. Separation from the Urine. (See Chapter 31.)
- 2. Melting Point. Determine the melting point of the hippuric acid prepared in the above experiment (see p. 793).
- 3. Formation of Nitrobenzene (Lücke's Reaction). To a little hippuric acid in a small porcelain dish add i to 2 ml. of concentrated HNO; and evaporate to dryness on a water bath. Transfer the residue to a dry test tube, apply heat, and note the odor of nitrobenzene (resembling that of oil of bitter almonds).
- 4. Sublimotion. Place a few crystals of hippuric acid in a dry test tube and apply heat. The crystals are reduced to an only fluid which solidifies in a crystalline mass upon cooling. When stronger heat is applied the liquid assumes a red color and finally yields a sublimate of benzoic acid and the odor of hydrocyanic acid.

OXALIC ACID

COOH

Ovalic acid is a constituent of normal urine, about 15 to 20 mg being channated in 24 hours. It separates out from neutral or alkaline urine

as the insoluble erystalline calcium oxalate, in the form of either dumbbells or octahedra, usually the latter (see Fig 230) Many urmary calcula consist largely of calcium ovalate When ingested ovalie acid is eliminated, at least in part, unchanged Since many common articles of diet, eg, asparagus, apples, cabbage, grapes, lettuce, rhubarb, spinach, tomatocs, etc, contain oxalates (or precursors) it seems probable that they are responsible for most of the oxalie acid found in the urine The ingestion of rhubarb leaves has caused oxalie acid poisoning. They contain about 12 per cent of the acid Spinach contains 08 to 09 per cent The eating of these vegetables tends to lessen the supply of calcium in the body Other green vegetables such as lettuce, celery, endive, asparagus, broccoli, Brussels sprouts, and cabbage contain very small amounts of oxalic acidgenerally less than 0 1 per cent There is also experimental evidence that part of the oxalic acid of the urine is formed within the organism in the course of protein and fat metabolism It has also been suggested that oxalic acid may arise from an incomplete combustion of carbohydrates especially under certain abnormal conditions. In this connection it 15 of interest that oxalic acid is one of the end products of in vitro oxidation of ascorbie acid Pathologically, urmary oxalic acid is increased in diabetes mellitus, in organic diseases of the liver, and in various other conditions which are accompanied by a derangement of the oxidative mechanism An abnormal increase in oxalic acid exerction is termed oxaluria and may be unaccompanied by any other apparent aymptom

Oxaluric acid (NH2 CO NH CO COOH) is occasionally found in traces in normal human urine. On hydrolysis it yields oxalic acid and urea

EXPERIMENT ON OXALIC ACID

Precipitation of Calcium Oxalate Place 200 to 250 ml of urine in a beaker add 5 ml of a saturated solution of calcium chloride, make the urine slightly acld with acelle acld, and sland the beaker in a cool place for 24 hours Examine the sediment under the microscope and compare the crystalline forms with those shown in Fig. 230, p. 855

CITRIC ACID

CH-COOH Ċ(OH)COOH ы.соон

Citric acid is a normal urinary constituent, the 24-hour excretion range ing from about 300 to 1500 mg Several factors have been noted to influ ence the excretion rate it is enhanced in alkaline urine, by estrogens, and with increased urmary excretion of calcium. Urmary citric acid excretion is reduced in acid urine and by androgens The effect of endogenous estrogens in augmenting urmary citric acid excretion is seen in

Ostberg Skand Arch Physiol 62 81 (1931)
 Shorr Bernheim and Taussky Science 95 606 (1942)

Shorr J Urology 51, 507 (1945)

the characteristic curve of the normal menstrual cycle, with the lowest excretion during menstruation followed by a mid-menstrual peal, and plateau and a return to the lower menstrual levels just before the next menstrual period. Since the citrate ion forms a soluble weakly ionized complex with calcium, an effort has been made to apply this augmenting effect of estrogens to the management of renal stones containing calcium. This application of estrogens is liable to be ineffective owing to the frequent association of urmary infections with organisms which destroy citric acid and is further limited in usefulness by its interference with menstrual cycles and with libido in the male. An enzyme, citrogenase, found in the kidney, heart, and liver can form citric acid by condensation of pyruvic and oxalacetic acids. A marked increase in urmary citric acid following hepatectomy suggests that the liver has a function in its intermediary metaholism.

NEUTRAL SULFUR COMPOUNDS

Under this head may be classed such substances as cystine, methyl mercaptan, ethyl sulfide, thoeyanates, taurine derivatives, etc. The sulfur content of the compounds just enumerated is generally termed unoxidized or neutral sulfur in order that it may not be confused with the aeid or oxidized sulfur which occurs in the morganic sulfuric acid and ethereal sulfurie acid forms, although this distinction is admittedly inexact. Ordinarily the neutral sulfur content of normal human urine is 5 to 25 per cent of the total sulfur content (see "Partition of Urinary Nitrogen and Sulfur," Chapter 31). The actual amount excreted may be 0.08 to 0.16 g. per day, calculated as S. Its origin is mainly endogenous. The excretion is fairly constant for any given individual in spite of dictary changes. In cystinuria, or in certain degenerative diseases, the amount is increased (See p. 168 for test for cystine and cysteine sulfur.)

ALLANTOIN

NH CH NH CO NH,

Allantom is found in the urine of practically all mammals including man In human urine it occurs in very small amounts (5 to 15 mg per day) whereas in all other mammals everyt anthropoid apes, and the Dalmatian coach dog, it is the principal end product of purine metabolism and may constitute 90 per cent or over of the total purine output Allantom is formed upon ovidation of urie acid by the enzy me uricase and the output is increased by the feeding of thymus or pancreas to lower animals. In the dog according to Mann and associates, the liver is the sole seat of urical edectruction, since after hepatectomy the urical edectruction, since after hepatectomy the urical edectruction, since after hepatectomy the urical edectruction, since after hepatectomy the urical edectruction, since after hepatectomy the urical edectruction is equivalent to the allantom output of the normal animal. When pure, allantom crystallizes in prisms (Fig. 221), and when impure in granules and knobs.

EXPERIMENTS ON ALLANTOIN

1 Separation from the Urme: Messager's Method. Precipitate the uring with baryta water Neutralize the filtrate carefully with dilute sulfurle add filter immediately, and evaporate the filtrate to incipient crystallization Completely precipitate this warm fluid with 95 per cent alcohol (reserve the complete) precipitate this warm fluid with 95 per cent alcohol (reserve the complete).



Tio 224 ALIANTOIN FROM CATS URINE
a, b, Forms in which it crystallized from
the urine a recrystallized allandor

precipitate), Decant or filter an precipitate the solution by ether Combine the ether and alcohol pre cipitates, and extract with color water or hot alcohol, aliantoin re mains undissolved. Bring the alian toin into solution in hot water and recrystallize

2 Preparation from Uric Acid Dissolve 4 g, of uric acid in 100 ml of water rendered alkaline with po tassium hydroxide Cool and care fully add 3 g of potassium per manganate Filter, immediately acidulate the filtrate with arctic acid, and allow it to stand in a cool place overnight Filter off the crystals and wash them with water Save the wash water and firrate

the urine c, recrystallized allanton

Save the wash water and fitrate unite them, and after concentrating to a small volume stand away for crystallization. Now combine all the experiments which follow

- 3 Microscopical Examination Examine the crystals made in Exp 2 and compare them with those shown in Fig. 224
- 4. Furfural Test (Schiff) Place a few crystals of aliantoin on a test tablet of in a porcelain dish and add it to 2 drops of a concentrated aqueous solution of furfural and it to 2 drops of concentrated hydrochiloric acid Observe the formation of a yellow color, which turns to a light purple if allowed to stand This test is given by urea but not by uric acid
- 5. Murexide Test Try this test according to the directions given on p. 796 Note that aliantoin fails to respond

AMINO ACIDS

Certain of these acids are always present in normal urine. The excretion of total amino and nitrogen by a normal adult averages 0.4 to 1.0 g per day or about 2 to 6 per cent of the total nitrogen. Tree amino and nitrogen (for methods of estimation, see Chapter 31) is considerably less than this and ordinarily constitutes 0.5 to 1 per cent of the total nitrogen. The

^{*} The urine of the dog after thamus pancreas or une acid feeding may be employed

amount may be largely increased in disorders associated with tissue waste, e g . typhoid, acidosis, pronounced atrophy of the liver, etc. After extirpa tion of the liver, urea formation crases and amino acids accumulate in the blood or are excreted in the urine if repal activity is maintained. For tests on amino acids, see Chapter 4

The availability of microhiological assay (see p 1061) and paper chromatography (see p 14) has made possible the detection and estima tion of the individual amino acids present in normal and pathological urine In one study on 18 normal male and female subjects on a normal diet, Woodson, et al 10a found most of the common amino acids present in the urine, in either the free state or in a combined form or both. The mean 24-hour excretion of free amino acids rauged from approximately 1 mg for aspartic acid to 188 mg for histidiue, for the combined form, from 2 mg for methionine to 315 mg for glutamic acid. Those amino acids excreted predominantly in the combined form included aspartic acid, glutamic acid, proline, valine, and isoleucine The nature of the combined form is not known, either peptides or conjugates are possibilitics No correlation could be established between the excretion of the various amino acids and differences in urine volume, total nitrogen, uric acid. creatinine, or ammonia content of the urines examined

Dent has made extensive studies of urmary excretion of amino acids. using the techniques of paper chromatography to which he has made valuable contributions 106 Investigations have been reported on the gen eral aminoaciduma characteristic of Fancom's syndrome, on the excessive excretion of cystine, lysine, and arginine in eystimuria, and on other

anomalies of amino acid metabolism

AROMATIC OXYACIDS

Two of the most important of the oxyacids are p-hydroxyphenylacetic acid, and p hydroxyphenylpropionic acid. They are products of the putre faction of protein material and tyrosine is an intermediate stage in their formation Both these acids for the most part pass unchanged into the urine, where they occur normally in very small amount The content may be increased in the same manner as the phenol content, in particular by acute phosphorus poisoning A fraction of the total aromatic oxyacid content of the urine is in combination with sulfurie acid, but the greater part is present in the form of salts of sodium and potassium

Levine11 has shown that the urine of premature infants fed cow's milk regularly contains certain aromatic oxyacids, such as p-hydroxyphenyl pyruvic acid, the product of the deamination of tyrosine Administration of vitamin C abolishes this defect. A similar observation has been noted by Scalock in the case of scorbutic guinea pigs (see discussion, p. 1043)

ive Woodson Hier Solomon and Bergeim J Bsel Chem. 172 613 (1948)
 ab Dent Biochem J 41 240 (1947) 43, 169 (1948) Biochem Soc Sympona 3 (Parts ton Chromotography) 34 (1959) Dent and Ro e Quart J Vied 20 205 (1941)
 ive Levine Science 39 (70 (1939) J Chin Irrest 22 351 (1943) See also Woolf and Edmunds Biochem. J 47 630 (1950)

Homogentisic acid or 2 5-dihydroxyphenylacetic acid,

is another important oxyacid sometimes present in the urine. Under the name glycosuric acid it was first isolated from the urine by Marshall, subsequently Baumann isolated it and determined its chemical constitution. It occurs in cases of alcaptonuria (a so-called "inborn error of metabolism") and has also been found in the urine of the scorbutic guinea pig A urine containing this oxyacid turns greenish brown from the surface downward when treated with a little sodium hydroxide or ammonia if the solution be stirred the color very soon becomes dark brown or even black. Homogentisic acid reduces alkaline copper solutions but not alkaline bismuth solutions.

Kynurence acid or γ hydroxy β -quinoline carboxylic acid is a product of the metabolism of tryptophan but it appears to be excreted only affect the ingestion of this amino acid in excess of normal requirements (see p. 1040)

Its presence after tryptophan administration in the urine of the dograt, rabbit hyens coyote wolf etc and absence from that of the evet cheetah, bear, raccoon etc (Jackson) may have significance in the zoological classification of species

EXPERIMENT ON KYNURENIC ACID

Isolation of Kynurenic Acid Aciddify the urne with hydrocbloric acid in the proportion 1.25 From this acid fluid both the uric acid and the kynurenic acid separate in the course of 24 to 48 hours. Filter off the combined crystal line deposit of the two acids, dissolve the kynurenic acid in dilute amount (uric acid is insolubic), and reprecipitate it with hydrochloric acid and into containing kynurenic acid be evaporated to dryness with hydrochic acid and potassium chlorate, are doise residue is obtained which becomes first brownish green and then emerald green on adding ammonia (gafe) is hynurenic acid may be quantitatively determined by Capaidi's method;

PROTEIN

The nubecula of normal urme has been shown by one investigator to consist of a mucoid containing 12 7 per cent of nitrogen and 2 3 per cent of

¹¹ Capalda Z physiol Chem 21 92 (1897) Berg J Biol Chem 91 513 (1931)

sulfur This substance evidently originates in the urinary passages. It is probably slightly soluble in the urine Some investigators believe that the material forming the oubscula of normal urine is nucleoprotein and not a mucin or mucoid. A discussion of nucleoprotein and related substances occurring in the urine under pathological conditions will be found in Chanter 29.

Normal urne contains a small amount of soluble protein—albumio, various enzymes (see below), etc., the amount is too slight to be detected by any but the most delicate procedures. The significance of pathological proteining is discussed in Chapter 29.

CARBOHYDRATES

Normal human unne cootains total reducing substances equivaleot to about 0 05 to 0 15 per cent of glucose. Fermentable sugars make up about one-tenth of this or about 0 01 per cent. Whether all of the fermeotable sugar is glucose is uncertain. Most of the sugar of normal unne is at any rate not glucose. The other sugars may include peotose, lactose, and altered carbobydrates formed by the baking of foods or through the action of bacteria in the intestinal tract. That glucose is not ordioarlly excreted in more than traces is further indicated by the failure of glucose logistion up to 1 g per kg of body weight to locrease the fermentable unne sugar. In various types of nephritis the excretion of fermentable sugar is increased somewhat. Lactose may be found in the union of pregnant women in appreciable amounts. In the rare conditions known as pentosuria and fructosuria these respective sugars may be present in readily detectable amount. Otherwise sugars are not cormally found in sufficient amounts to give the ordinary sugar tests. Pathological glucosuria is discussed in Chapter 29.

ENZYMES

Various types of enzymes produced within the organism are excreted in both the feces and the urine. In this connection it is interesting to note that pepsin, trypsin, lipase, and an amylase have been positively identified in the urioe. The amylase may be much increased in panereatic disease.

VITAMINS

The normal path of elimination of the fat-soluble vitamins is the intestinal tract. The water-soluble vitamins thiamine, riboflavin, ascorbie acid, and others, are evereted in the urine under normal conditions of vitamin nutrition, the amounts of these vitamins in the 24-hour everetion being directly proportional to the intake. Nicotinamide is normally eliminated in the form of the metabolite N*-metbylineotiuamide. The urinary output of the water-soluble vitamins under controlled conditions forms the basis of clinical tests for vitamin deficiencies, since in such states the tissues are unsaturated and tend to retain the test doses of vitamins. In a study on seven health, males on a cormal dietary regime¹⁴ the following ranges of values were found for daily vitamin mixike and output

showing ranges of varies were found for daily vicanini incake and our

¹¹ Dodds Brit J Expil Path 3, 133 (1922)
11 Denko et al | 1rch. Biochem 10 33 (1946) 11 109 (1946)

Chap 28

Vitamin Thiamine mg. Riboflavin, mg Viscen, mg. Vislethylin otinamide, mg. Pantothenie acid, mg. Pyridoxine, mg. Folic acid, #g. Biotin, #g Palminobenzoic acid, #g.	Intale	Outp	Areray Pzerelion	
	Intare	Urine	Feces	Per Cen
	1 74- 1 98 12 4 -20 9 - 4 13- 5 30	0 144-0 323 0 543-0 913 1 13-1 39 2 7 - 4 4 2 68-3 46 0 57-0 6J 2 94-4 99 27 8-35 6 131-198	0 823-1 313	31 112

^{*} Per cent excretion = 100 (Urmary + Lecal output)/Intake

The high excretion rates of the last three vitamins (particularly via the fecal route), is a reflection of the extent of synthesis by intestinal flora. The figures in this table do not represent the entire exerction of nicotime acid and pyridoxine since their metabolites, V—methylnicotinamide and 4-py ridoxic acid, respectively, were not included

VOLATILE FATTY ACIDS

Acette hutyric, and formic acids have been found under normal conditions in the urine of man and of certain carmivors as well as in the urine of herbityria. Normally they arise principally from the fermentation of car hohydrates and the putrefaction of proteins. The acids containing the few est carbon atoms (formic and acette) are found to be present in large percentage than those which contain a larger number of such atoms. The volatile fatty acids occur in normal urine in traces, the total output for 2 hours according to older investigators varying from 0.008 g. to 0.05 formic acid exerction is increased in methyl alcohol poisoning.

LACTIC ACID

Lactic acid is supposed to pass into the urine when the supply of orgen in the organism is diminished through any cause, e.g., in pneumonicalmpsia, acute yellow atrophy of the liver, carbon monoxide poisonin acute phosphorus poisoning or epileptic attacks. This acid has also be found in the urine of bealthy persons following the physical exercimendent to prolonged marching Liljestrand and Wilson found the outy of lactic acid to vary from 140 mg to 1370 mg after a few minutes stremuous physical exercise e.g. stair running (See experiment, Chapi 33) Lactic acid has been detected in the urine of birds after the remotor the liver.

PHOSPHORYLATED COMPOUNDS

Phosphorus in organic combination has been found in the urine in substances as glycerophosphore and which may arise from the decoposition of lecithin, and phosphocarnic acid. It is claimed that out to

average about 25 per cent of the total phosphorus elimination is in organic combination

PIGMENTS

There are several pigments normally present in human urine, of which the most important is urochrome. Small amounts of urobilin, urocrythrin, and certain porphyrins (uroporphyrin and coproporphyrin) are also present.

Urochrome Urochrome is the principal pigment of normal urine. Its chemical unture is not definitely established. It may be a compound of urobilin and urohilinogen with a peptide substance. It is a product of endogenous metabohsm and is fairly constant in amount from day to day in the urine of normal individuals.

Urohilm. Urobilu is normally present in too small amount to give any appreciable color to the urine. It exists chiefly in the reduced form as the colorless urobilinogen which upon oxidation gives urobilin. (See Chapter 18.) It is derived from bile pigment by bacterial action in the bowel and most of it is excreted in the feces where the corresponding and probably identical compounds are known as stereobilinogen and stereobilin, respectively. Of the small amount absorbed the larger part is excreted in the bile.

PURINE BASES

The purine bases found in human urine are adenine, epiguanine (7 methylguanine), guanine, vanthine, heteroxanthine (7-methylxanthue), hypoxanthine, paraxauthine (1,7 dimetby kanthine), and 1-methylxanthine The main bulk of the purine base content of the uriue is made up of 7-methy lanthine, 1,7-dimethylanthine and 1 methyl xanthine, which are derived for the most part from the caffeine, theobromue, and theophylhne of the food The total purme base content is made up of the products of two distinct forms of metabolism, i.e. metabohism of ingested nucleoproteins and purines and metabolism of tissue nuclear material Purine bases resulting from the first form of metabolism are said to be of exogenous origin, whereas those resulting from the second form of metabolism are said to be of endogenous origin. The daily output of purme bases by the time is extremely small and varies greatly with the individual (16 to 60 mg) The output is increased after the ingestion of nuclear material as well as after the increased destruction of leukocytes A well marked increase accompanies leukemia. The output of purino bases by the urine is increased as a result of a ray treatment. The purine bases form a higher percentage of the total purme excretion in the case of the monkey, sheep, and goat than in man

EXPERIMENT ON PURINE BASES

Formation of the Silver Salts Add an excess of magnesia mixture. to 25 ml of urine Filter off the precipitate and add ammoniacal silver solution to the filtrate, A precipitate composed of the silver salts of the various purine bases is produced. The purine bases may be determined quantitatively by Kruger and Schmidt s method or Welker's method (see Chapter 31).

¹⁴ See Appendix.

INORGANIC PHYSIOLOGICAL CONSTITUENTS

AMMONIA

Next to urca, ammonia is quantitatively the most important of the nitrogenous end products of protein metabolism Ordinarily about 23 to 45 per cent of the total nitrogen of the urine is eliminated as ammonia and on the average this would be about 07 g per day The significance of the ammonia content of the urine appears to he primarily if not entirely concerned with the mechanisms of acid-hase balance in the hody (see Chapter 24) If ammonia is fed in the form of oxidizable organic salts, such as ammonium acetate, ammonium lactate, etc., no extra ammonia appears in the urine The organic portion of the salt is oxidized, and the ammonia portion is converted into urea and excreted in this form If, however, a nonoxidizable salt such as ammonium chloride is administered while the ammonia portion is probably also converted into urea for excretion (since there is no reason to suppose that the metabolism of the ammonium ion differs with different salts) the excretion of the extra chloride ions requires the simultaneous exerction of an equivalent amount of base This hase must be either sodium, potassium, or ammonium 1003 themselves, since these are the only forms of hase available to the kidney In the presence of abundant sodium or potassium (fixed hase), the extra chloride is excreted largely as the sodium or potassium salt and no disturhance of acid hase balance results If, however, the supply of fixed hase is limited, its excretion results in the development of an acidosis due to the depletion of hody hase, in fact, the administration of a large dose of ammonium chloride is used clinically for the production of an acidosis The kidney therefore synthesizes ammonia (volatile hase) to the limit of its capacity under these conditions to conserve body base, the extra am monia is excreted in the urine and the urinary ammonia content rises Excretion of acid phosphate also occurs here (see p 818) That the in gested ammonia of the ammonium chloride plays no important part in the increased exerction of ammonia is shown by the fact that exactly the same condition occurs if hydrochloric acid is administered instead of ammonium chloride there is a rise of urinary ammonia accompanying the increased excretion of chloride Acid forming foods (see Chapter 34) also increase the ammonia output whereas the administration of alkalies or of base-forming foods decreases the excretion of ammonia Copious water drinking increases the ammonia output This fact has been interpreted as indicating a stimulation of the gastric secretion

The acids formed during the process of protein destruction within the body (i.e. sulfurn acid formed by oxidation of the sulfur of methionine and cystine) have an influence upon the excretion of ammonia similar to that exerted by acids which have been administered. Therefore a pathological increase in the output of ammonia is observed in such diseases as are accompanied by an increased and imperfect protein metaholism Likuwise in diabetes melhitus where the excretion of excessive amonits of acctoactic acid and 4-bydroxy butyre acid as their salts also tends to deprive the body of fixed base increased excretion of ammonia occurs.

.i

The kidney is the source of urinary ammonia. This is indicated by the fact that following extirpation of the kidneys the blood ammonia does not increase. On removal of the liver blood ammonia increases but not urinary ammonia. This indicates that the two have not the same origin. The blood ammonia appears to arise from the tissues, especially the muscles (probahly from the deamination of adenylic acid) and to he converted so readily to urea that the hlood ammonia remains too low to account for any appreciable part of the urinary ammonia. Ammonia is apparently formed by the cells of the tubules of the kidney. The precursor is believed to he an amide-nitrogen compound, probably glutamine, in blood plasma (Van Slyke) from which the enzyme glutaminase forms ammonia and glutamic acid. Deamination of amino acids by kidney tissue may also account for a small portion, possibly through the intermediate formation of glutamine. A decrease in the pH of the blood or of the kidney tissue seems to speed up the ammonia-producing mechanism.

The quantitative determination of ammonia must be made upon fresh or properly preserved urine, since upon standing normal urine will

undergo ammoniacal fermentation (see p. 784).

EXPERIMENT ON AMMONIA

(See Exp. 2 under "Experiments on Phosphates," p. 819.)

SULFATES

Sulfur in combination is excreted in two forms in the urine: first, as unoxidized, loosely combined or neutral sulfur, and second, as oxidized or acid sulfur. The excretiou of neutral sulfur has already been discussed (p. 807). The oxidized sulfur is eliminated chiefly in the form of the inorganic sulfate ion; a relatively small amount occurs in the form of ethereal sulfate, i.e., salts of sulfuric acid in combination with such aromatic substances as phenol, indole, skatole, cresol, pyrocatechol, and hydroquinol. This latter form of sulfurie acid is sometimes called conjugate sulfuric acid. The greater part of the total sulfur is climinated in the oxidized form, but the absolute percentage of sulfur excreted in the various forms depends upon the total quantity of sulfur present; i.e., there is no definite ratio between the three forms of sulfur which will apply under all conditions. The preformed or inorganic sulfurie acid may he precipitated directly from acidified urine with BaCl, whereas the ethereal sulfurie acid must undergo a preliminary hydrolysis hefore it can be so precipitated.

The sulfuric acid excreted in the urine arises principally from the oxidation of the sulfur of protein within the body; a relatively small amount is due to ingested sulfates. Under normal conditions about 1.0 g. of total S is eliminated daily, about 75 to 95 per cent of this being in the form of sulfates. About 90 per cent of this sulfate exerction is in the form of inorganic sulfate and 10 per cent as ethereal sulfates and neutral sulfur.

The sulfate ion is exercted with greater difficulty than any other inorganic radical ordinarily present in normal blood. A retention of 30 times the normal blood value (0.9 to 1.1 mg, of S per 100 ml, of whole blood) has been observed Sulfate ions are not readily absorbed by the tissues even when present in high concentration in the blood

EXPERIMENTS ON SULFATES

- 1 Detection of Inorganic Sulfuric Acid Place about 10 ml of urine in a test tube, acidify with acetic acid, and add some barlum chloride solution A white precipitate of barium sulfate forms
- 2 Detection of Ethereal Sulfuric Acid Filter off the barium sulfate precipitate formed in the above experiment, add 1 ml of hydrochloric acid and a little barium chloride solution to the filtrate and heat the mixture to boiling for 1 to 2 minutes Note the appearance of a turbidity due to the presence of sulfuric acid which has been hydrolyzed from the ethereal sulfates and bas reacted with BaCl; to form BaSO.
- 3 Detection of Unoxidized or Neutral Suffur Place about 10 ml of urine in a test tube, introduce a small piece of zinc, add sufficient hydrochloric acid to cause a gentle evolution of hydrogen, and over the mouth of the tube place a filter paper saturated with lead acetate solution in a short time the portion of the paper in contact with the vapors within the test tube becomes blackened due to the formation of lead sulfide. The nascent hydrogen has reacted with the loosely combined or near the state of the

this gas coming in contact with the lead acetate paper has caused the production of the black lead sulfade Sulfur in the form of inorganic or ethereal sulfuric acid does not respond to this test (For discussion of neutral sulfur compounds, see p 807)



4 Calcrum Sulfate Crystals Place 10 ml of urms in a test tube, add 10 drops of comparts of urms in a test tube, and 10 drops of urms in a test tube, and allow the tube to stand until crystals form Examina the calcrum sulfate crystals under the

microscope and compare them with those shown in Fig. 225

CHLORIDES

Next to urea, the various chlorides constitute the chief solid constituent of the urine. The excretion of chloride is dependent in great part, upon the nature of the diet hut on the average the daily output is about 10 to 15 g expressed as sodium chloride. Conditions which favor excessive perspiration, such as stremous athletic or occupational activity, especially in a hot environment cause a diminution in urinary output of chlorides, the chloride leaving the body by other channels. Muscular eramps may result when there is an excessive loss of sodium cbloride from the body. To objuste this condition workers in a high temperature environment or persons indulging in very fatiguing muscular efforts, such as athletes in prolonged competition make use of saft tablets. Because of their solubility chlorides are never found in the urinary sediment.

The amount of chlorides exerted in the urner is related primarily to the chloride content of the food ingested. In cases of actual fasting the chloride content of the urine may be decreased to a slight trace which is derived from the body fluids and tissues Under these conditions, however, au examination of the blood of the fasting subject will show the content of chloride in this fluid to be approximately normal. This forms a very striking example of the care nature takes to maintain the normal composition of the blood. There is a limit to the power of the body to maintain this equilibrium however, and if the fasting organism be subjected to the influence of diuretics for a time, a point is reached where the normal composition of the blood can no longer be maintained and a gradual decrease in its chloride content occurs which finally results in death Since the excreted chloride must carry base (e.g., sodium) along with it, death results not so much from the loss of chloride alone as from the mability of the organism to maintain a normal osmotic pressure in its body fluids, which is a major function of the sodium and chloride ions Potassium cannot take the place of sodium in this respect thus the administration of notassium chloride under these conditions is of no value whatever

Pathologically, the exerction of chlorides may be decreased in some fevers chronic nephritis, fasting, diarrhea, certain stomach disorders, and after extensive burns. Any condition accompanied by the formation of an exudate (e.g., pneumoua) will cause a diminished chloride output. In convalescence and with resolution of the exudate the chloride exerction 1888 again.

EXPERIMENTS ON CHLORIDES

Detection of Chlorides in Urine Place about 5 ml of urine in a test tube, render it acid with nitric acid, and add a few drops of a solution of silver nitrate A white precipitate, due to the formation of silver chloride, is produced This precipitate is soluble in ammonium hydroxide

PHOSPHATES

Of the various inorganic amons of urine, phosphate is ordinarily second only to chloride in amount present. The exerction of phosphate is extremely variable, depending in large measure upon the diet, but on the average the total output for 24 hours is about 1 I g expressed as P The bulk of this is in the form of morganic phosphate, the organic phos phorus of the urme constituting only about 1 to 4 per cent of the total phosphorus content The greater part of the morganic phosphate arises from the ingested food, either from phosphate already present as such or more especially from the metabolism of compounds containing phosphorus in organic combination, such as phosphoproteins nucleoproteins and nucleotides and phospholipides The various phosphorus-containing compounds of the hody also contribute to the total output of this element The phosphate content of the urme also depends to a considerable extent on conditions within the intestinal lumen, an increased alkalimity together with the presence of substruces like calcium and magnesium which form insoluble phosphates tends to increase the proportion of phosphate excreted in the feces at the expense of the urmary pho phate content

The phosphite ion is found in the urine in two forms, the acid phosphate or monohasic ion, H-PO, and the dibasic ion, HPO, The ratio of

these two ions determines in large measure the pH of the urine, since they constitute the major buffer system ordinarily present. In blood and in the glomerular ultrafiltrate of the kidney, these ions are present in a ratio corresponding to about 80 per cent basic phosphate and 20 per cent acid phosphate If this ratio prevails in the urine, it will have the same pH as the blood, or approximately 7.4 Increased acidity of the urine is due to an increase in the amount of the acid phosphate relative to basic phosphate, at pH 6 6, which is approximately the pH of average urine, about 60 per cent of the total morganie phosphate is in the and form and 10 per cent appears as basic phosphate. The best avadable evidence indicates that the ability of the kidney to excrete a varying fraction of its total phosphate in either the acid or hasie forms is an important part of the mechanism for the regulation of acid-base balance in the body, actually, the excretion of one equivalent of acid phosphate instead of hasic phosphate corresponds to the excretion of one equivalent of hydrogen ion itself, and the simultaneous retention of one equivalent of base, as inspection of the formulas of these two salts will show. In combating acidosis, this mechanism appears to be of importance second only to the ability of the kidneys to replace sodium or potassium by ammonium Available evidence indicates that the change from acid to hasie phosphato and vice versa is brought about in the renal tubules, possibly hy secretion of hydrogen ions (Pitts), since the total phosphate excretion does not vary with changes in the relative amounts of the acid and basic forms excreted

If urine containing phosphate is rendered sufficiently alkaline, the so-called "alkaline earth" elements calcium and magnesium will precipitate, to the extent of their presence, as insoluble calcium and mag nesium phosphates. This fraction of the total phosphate of urine was formerly called the "earthy phosphate" fraction as contrasted to tho "alkaline phosphate" fraction composed of the soluble phosphates of sodium, potassium, and ammonium It is doubtful whether this fractionation has any significance other than to indicate the amount of calcium and magnesium present relative to the total phosphate, and it should be abandoned

The so-called "phosphaturia" the appearance of a copious crystalline precipitate which on examination proves to be magnesium ammonium phosphate ("triple phosphate"), ordinarily represents a decreased acidity of the urine and not an increased phosphate content. Such conditions may, however, be of aignificance in connection with a possible tendency to the formation of phosphate calculi Measures designed to acidify the urine, and to decrease its phosphate content, have been of value in the treatment of this condition

Pathologically the excretion of phosphoric acid is increased in such discases of the bones as diffuse periostosis, osteomalacia, and rickets, according to some investigators in the early stages of pulmonary tuberculosis, in acute yellow atrophy of the liver, in diseases which are accompanied by an extensive decomposition of nervous tissue, and after sleep induced by potassium bromide or chloral hydrate (Mendel) It is also increased after copious water drinking 1 decrease in the excretion of phosphates is at times noted in febrile affections, such as the acute infectious diseases, in pregnancy, in the period during which the fetal bones are forming, and in diseases of the Lidneys, because of nonelimination

EXPERIMENTS ON PHOSPHATES

1. Formation of "Triple Phosphate." Place some urine in a beaker, render it slightly aikaline with ammonium hydroxide, add a small amount of magnesium sulfate solution, and allow the heaker to stand in a cool place overnight. Crystals of ammonium magnesium phosphate, "triple phosphate," form under these conditions. Examine the crystaline sediment under the microscape and compare the forms of the crystals with those shown in Fig. 226. If possible, examine the crystals from a freshly passed specimen of cloudy urine, or from a sample of urine which on standing for a short while after

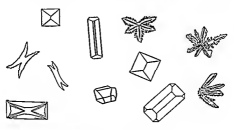


FIG 226 TRIPLE PHOSPHATE (OGDEN)

voiding has become cloudy. How do they compare with those obtained in this experiment?

- 2. Ammoniacal Fermentation. Stand some urine aside in a beaker for several days. Ammoniacal fermentation will develop and "triple phosphate" crystals will form
- Examine the sediment under the microscope and compare the crystais with those shown in Fig. 226.

b. Hold a glass rod dipped in concentrated hydrochioric acid near the surface of the urine. Note the fumes of ammonium chioride.

c. insert a strip of red litmus paper in the urine. Permit the paper to dry. Note the gradual restoration of red coior, due to voiatilization of ammonia (voiatile aikaii). Run a control test using 0 5 per cent Na-CO, (fixed alkaii).

SODIUM AND POTASSIUM

Sodium and potassium ions are always present in the urine. The amount of potassium excreted in 24 hours by an adult, subsisting upon a mixed diet, is on the average 1 to 3 g, whereas the amount of sodium under the same conditions is ordinarily 3 to 5 g. The ratio of K to Na is generally about 3 5. The absolute quantity of these elements exercted depends, of course, in large measure upon the nature of the diet. Because of the noningestion of NaCl and the accompanying destruction of potas-

sium containing body tissues, during fasting the urine contains more notassium than sodium

Pathologically the output of potassium, in its relation to sodium, may be increased during fever, following the crisis, however, the output of this element may be decreased. It may also be increased in conditions associated with acidosis. In Addison's disease there is usually a relative retention of potassium and an increased excretion of sodium, the administration of cortical hormone preparations restores the normal relationship.

CALCIUM AND MAGNESIUM

The daily output of calcium in the urine, which depends principally upon the nature of the diet, is on the average about 0 1 to 0 3 g per day The percentage of calcium present in the urine at any one time (10 to 40 per cent of total calcium output) forms no dependable index as to the absorption of this element, since it may be again excreted into the audity or alkalinity of the intestinal lumen, and the presence of substances such as phosphate and fatty acids which form insoluble calcium salts, may determine to a considerable extent the relative output of calcium in the fixes and the urine Acidity promotes calcium absorption, alkalinity retards it. It is therefore impossible to draw any satisfactory conclusions regarding the excretion of calcium unless accurate analytical data from both the fexes and the urine are obtained.

Myers and Fine¹⁶ have reported data showing a comparison of the hidney and infestine as exerctory routes for various morganic constituents. Their findings in this connection are summarized in the following table

\umber	Mossiure Con tent of Feces (Per Cent)	Fecal Output in Per Cent of Total in 1 rine and Feces							
of Cases		H _z O	`	5	CI	P	Ca	Mg	К
5 9	76 84	6	10 13	10	3 9	36 33	90	72 68	18 27

The average findings in five cases with well-formed stools, 74 to 79 per cent moisture, and those with diarrheal stools, 79 to 89 per cent moisture, have been grouped separately in the table it is not behaved that the findings differed especially from the normal, except in that group of cases which suffered from intestinal diarrhea

Very little is known positively regarding the actual course of the excretion of calcium under pathological conditions. An excess is found in some diseases of the bones e.g., osteomalacia. In others, as in rickets, the unnary exerction may be very low.

The daily excretion of magnesium by way of the urine usually amounts to between 0.05 and 0.2 g. The amount depends mainly on the diet About 65 per cent or more of the exercted magnesium is usually eliminated by

¹⁴ Myers and Fine I roc Soc Expd Biol Med 16 73 (1919)

the feces; the remainder passes out in the kidneys. There may be a retention of magnesium in certain bone disorders accompanying a loss of calcium, for example, in osteomalacia. Thus the exerction of calcium and magnesium do not necessarily run parallel.

EXPERIMENT ON CALCIUM

Sulkowitch Test. This is a qualitative test for calcium in the urine. It has applications in parathyroid derangement, urinary calculi, infantile tetany, severe nephritis, etc.

Procedure. Place the patient on a diet containing sufficient calcium for his needs, and collect a 24-hour specimen of urine. To 5 ml. of the mixed urine sample add 2 ml. of the Sulkowitch reagent? dropwise. If no precipitate forms immediately, mix the contents of the tube thoroughly and allow to stand,

Interpretation. In case there is no precipitate there is of course no calcium in the urine. From this finding it is concluded that the blood serim probably contains not more than 5 to 75 mg per 100 ml If the precipitate forms as a fine white cloud, the calcium content of the scrim is considered to fall within the normal range of 9 to 11 mg per 100 ml. However if a heavy, milklike precipitate forms, there is danger of hypercalcemia. In routine examinations, the precipitates may be graded 1, 2, 3 or 4.

CARBONATES

Carbonate, in the form of the brearbonate ion, HCO₇, generally occurs in small amount in the neutral or alkalioe urine of man and carnivora, whereas much larger quautities are ordinarily present in the consistently alkaline urine of herbivora. The alkaline reaction of the urine of herbivora is due in great measure to the presence of bicarbonate. The carbonates of the alkaline earths are sometimes found in amorphous urinary sediments from quite alkaline urine.

IRON

Iron is present in small amount in normal urine. It probably occurs partly in inorganic and partly in organic combination. The iron contained in utinary pigments or chromogens is in organic combination. According to different investigators the iron content of normal urine probably averages not more than 1 to 2 mg. per day. Mitchell and Hamilton¹⁸ report that sweat contains 1-2 mg. of iron per liter. After splenectomy there is an increased loss of iron from the body particularly by way of the feces.

the presence of iron. (b) To the second part of the solution add a little potassium ferrocyanide solution; a precipitate of prussian blue forms upon standine.

FLUORIDES, NITRATES, AND SILICATES

These substances are all found in traces in human urine under normal conditions. Nitrates are undoubtedly introduced into the organism in the water and ingested food. The average excretion of nitrates is about 0.5 g. per day, the output being the largest upon a vegetable diet and smallest upon a meat diet. Nitrites are found only in urine which is undergoing decomposition and are formed from nitrates in the course of ammoniacal formentation.

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Urine: Pathological Constituents

Many of the substances considered in this chapter as pathological constituents of urine are present in small amount in normal urine. Hence their pathological significance may be more a question of the amount present than of their actual presence in or absence from urine. It is generally true, however, that the usual qualitative tests for these substances are of such sensitivity as to yield an essentially negative result when applied to normal urine, but to respond readily when unusual amounts are present. In this connection it is well to remember that a single specimen of urine may be sufficiently concentrated or otherwise influenced hy dietary or other factors as to yield a positive result with a particular test, the significance of which disappears, however, when considered in the light of the total 24-hour excretion Whenever possible, therefore, unless it is desired otherwise for special purposes, it is recommended that urine tests be carried out on a portion of the well-mixed and properly preserved 24-hour urine before interpreting results; if this is not feasible, the overnight sample collected before breakfast should be used.

GLUCOSE

Traces of this sugar may occur in normal urine, but the amount is ordinarily too small to be demonstrable by the common qualitative tests (see also Chapter 31). The presence of readily detectable amounts of glucose in urine is known as glucosuria; the term glycosuria is frequently used, hut this expression more correctly refers to the presence of any sugar, not necessarily glucose, in the urine. Glucosuria may he either benign or pathological, and it is important to distinguish hetween these two types. Renal diahetes, in which the kidney threshold for glucose is below normal hut the blood sugar level is normal, is an example of a henign glucosuria, as are the glucosurias associated with alimentary and emotional hyperglycemias. Pathological glucosurias include chiefly those of diahetes mellitus and other endocrine disorders, in which there is a marked elevation of the blood sugar and usually an increased volume of urine (polyuria). The glucose content of the urine in diabetes mellitus may he as high as 10 per cent or more, but is usually around 3 to 5 per cent. The urine may be light in color and have a high specific gravity.

TESTS FOR GLUCOSE

The various tests for glucose in the urine which are embraced in the experiments given herewith are based upon one of the following properties

of this sugar, as discussed in Chapter 2 (1) Its power to reduce the ions of certain metals in alkaline solution, (2) its power to rotate the plane of polarized light (3) its power to form a crystalline osazone with phenylhydrazine, and (4) its ability to ferment with ordinary yeast

None of these tests by itself is specific for glucose Positive evidence that the sugar is glucose may be obtained by demonstrating that the reducing power disappears after yeast fermentation and that a typical glucosazone is obtained in the absence of a positive test for fructose or mannose (see below) Quantitative measurement of optical rotation in relation to

total reducing power is also of value

The official test for glucose adopted by the Association of Life Insurance Medical Directors of America may be found in Chapter 31

1. Phenylhydrazine Reaction. Yellow crystalline compounds called osazones are formed from certain sugars by reaction with phenylby drazine, in general each individual sugar giving rise to an osazone of a definite

crystalline form which is typical for that sugar

As applied to urine, however, it is frequently difficult to obtain characteristic crystals, even for glucose, since crystal form and growth may be influenced by other substances present Better results are usually obtained if the urino is clarified first, as described below Results are more significant if they are positive (i.e., if characteristic crystals are obtained) than if they are negative, but in any event the test should be regarded as a guide or confirmation to be accompanied by more specific tests Identification of the individual sugar osazones by their melting points is of little value

In this connection it is important to remember that of the simple sugars of interest in physiological chemistry glucose fruetose and mannose give the same esazone Tructose may be ruled out by the absence of a positive Schvanoff test (see p 848) the presence of mannose will be indicated by the formation of a colorless crystalline hydrazone on treatment with phenylhydrazine in the cold, prior to the heating which produces the osazone

Procedure To a small amount of phenylhydrazine mixture1 (about one half inch in a small test tube), add 5 ml of the urine, shake well, and heat on a boiling water bath for one half to three quarters of an hour Allow the tube to cool slowly (not under the tap) and examine the crystals micro scopically (see Plate II, opposite p 63) If the solution has become too con centrated in the boiling process it will be light red in color and no crystals will separate until it is diluted with water

in case doubtful results are obtained by this test owing to the presence of interfering substances, the urine should be clarified and the test repeated To ciarify the urine introduce 10 ml into a test tube, add 1 g of pure blood charcoal, heat to boiling and allow to stand with occasional shaking for five minutes, then filter Lee the filtrate in the test

2. Reduction Tests. It is to their potential aldehyde or ketone structure that many sugars owe the property of readily reducing alkaline solutions of metals like copper, bismuth mercury, and iron, they also

¹ See Appendix.

possess the property of reducing ammoniacal silver solutions with the separation of metallic silver. Upon this property of reduction the most widely used tests for sugars are based. A positive reduction test is not specific for glucose, or even for reducing sugars in general, non-sugar-reducing material is present in traces in normal urine and may be sufficiently augmented in amount, particularly in concentrated urines, to give a positive test. Other factors which may influence interpretation of results are discussed below. The chemistry of the various reduction tests and reagents is discussed in Chapter 2.

a Fehling's Test To ahout 1 ml of Fehling's solution in a test tube add about 4 ml of water, and holl 'This is done to determine whether the solution will of itself cause the formation of a precipitate of brownish red cuprous oxide 1f such a precipitate forms, the Fehling's solution must not be used Add urine to the hot Fehling's solution, a few drops at a time, and heat the mixture to holling after each addition (never add more urine than the original volume of Fehling's solution) The production of yellow or brownish red cuprous oxide indicates that reduction has taken place. The yellow precipitate is more likely to occur if the urine is added rapidly and in large amount, whereas with a less rapid addition of smaller amounts of urine the brownish red precipitate is generally formed. The differences in color of the exprous oxide precipitates under difference conditions are apparently due to differences in the size of the particles, the more finely divided precipitates having a yellow color, while the coarser ones are red. In the presence of protective colloids substances the yellow precipitates is usually formed.

Fhis classical test is not entirely rehable when used to detect sugar in the urine, and has been largely replaced by Benedict's test (see below) Such compounds as conjugate glucuronates, uric acid, nucleoprotein, and homogentisic acid, when present in sufficient amount, may produce a result similar to that produced by sugar Phosphates of the alkaline earths may be precipitated by the alkalin of the Fehling's solution as a grayish white precipitate which should not be mistaken for the cuprous oude Cupric hydroude may also be reduced to cuprous oude and this in turn may form a soluble complex with creatinine, a normal urinary constituent. This will give the urine under examination a greenish tringe and may obscure the sugar reaction even when a considerable amount of sugar is present.

Conjugate glucuronates are present in normal urine in small amount (see p. 842) and are increased in amount after the nigestion of such substances as chloral hydrate, camphor, menthol, thymol, antipyrine, phenol, etc. The chloral hydrate is excreted in the urine as trichloroethyliglucuronate. This compound reduces Felhing's solution and is levorotatory, whereas glucose also reduces but is destrorotatory. Therefore by means

See Appendix

More distinct Tehling a solution should be used in testing urines containing small amounts of sugar. In case of urines containing a high concentration of sugar it may sometimes be learnable to use a Lirger volume of 1 tl ling a solution.

In case doubtful results are channel by H is test owing to the presence of interfering substances the urice should be clarified with charced as described above and the test repeated

of a polariscopic test a "chloral urine" may be differentiated from a "sugar

urine " In testing urine preserved by chloroform a positive test may he obtained in the absence of sugar This is due to the fact that the hot alkali produces a reducing substance from the chloroform

Ammonium salts also interfere with Fehling's test preventing the precipitation of cuprous oxide If they are present in excess, the urine should be made alkaline with strong sodium carbonate, and boiled (or, better, aerated with a vigorous stream of air) to decompose and liberate the ammonta

b Benedict's Test This is the most satisfactory of the copper reduction tests, and in laboratory practice has largely replaced Fehling's The following is the procedure for the detection of glucose in urine To 5 ml of the reagent's in a test tube add exactly 8 drops of the urine to be examined The fluid is then holled vigorously for from one to two minutes and then allowed to cool spontaneously (Do not hasten cooling by immersion in cold water) The test may also be carried out by heating for five minutes in a boiling water hath, removing, and allowing to cool in the air This procedure is recom mended for serial tests If a water hath is used, it is important that the urine and reagent he thoroughly mixed before placing in the water hath. In the presence of glucose the entire body of the solution will he filled with a col loldal precipitate, which may be green, yellow, or red in color, depending upon the amount of sugar present In the presence of over 0 2 to 0 3 per cent of glucose, the precipitate will form quickly If no glucose is present, the solution will either remain perfectly clear, or will show a very faint turbidity. due to precipitated urates

Even very small quantities of gluco-e in urine (0 1 per cent) yield precipitates of surprising bulk with this reagent and the positive reaction for glucose is the filling of the entire body of the solution with a precipitate, so that the solution becomes opaque Since amount rather than color of the precipitate is made the basis of this test it may be applied, even for the detection of small quantities of gluco.e, as readily in artificial light as in daylight Chloroform does not interfere with this test nor does uric acid or creatinine interfere to such an extent as in the case of Fehling's test It is common clinical practice to run this test under consistently uniform conditions (same amount of urine heating time etc.) and to indicate the intensity of the reaction by arbitrary signs (+, ++, +++, etc) as an index not only of the presence of sugar but also of its approxi mate relative concentration *

c Bismuth Reduction Test (hylander) To 5 ml of urine in a test tube add one tenth its volume of 'ylander a reagent' and heat for 5 minutes in 2

bee Appendix.

A sample modification of the copper reduction test for urine augar is available com mercially in the form of tablets containing copper sullate sodium hydroxide and citric acid (Chrutest tablets obtainable from the Ames Co Inc. Elkhart Indiana) In use a little urine is diluted with 2 volumes of water and a tablet added to the mixture. Apply caise n of external heat is unnecessary. After the mixture has stood for a few moments, the 1) pical appearance of a positive copper reducts in test is obtained if the urine contains reducing sugar

boiling water bath. The mixture will darken if reducing sugar is present, and upon standing for a few moments a black color will appear.

This color is due to the precipitation of bismuth.* If the test is made on urine containing albumin this must be removed, by hoiling and filtering, before applying the test, since with albumin a similar change of color is produced. Glucose when present to the extent of 0.08 per cent or even less may be easily detected by this reaction. Uric acid, creatinine, and homogentisic acid which interfere with the Fehling test, do not interfere with the Nylander reaction.

Urine rich in indican, uroerythrin, urochrome, or porphyrins, as well as urine excreted after the ingestion of large amounts of certain drugs, may give a darkening of the Nylander's reagent similar to that of a true sugar reaction. It has been claimed that the urine after the administration of urotropin will reduce the Nylander reagent

A positive hismuth reduction test is probably due to the following reactions:

$$Bi(OH)_2NO_2 + KOH \rightarrow Bi(OH)_3 + KNO_3$$

 $2Bi(OH)_3 - 3O \rightarrow 2Bi + 3H_3O$

Before testing the urine, Bohmansson treats 10 ml with 2 ml of 25 per cent hydrochloric acid and 4 to 5 g of boneblack. This mixture is shaken one minute, then filtered, and the neutrabzed filtrate tested by Nylander's reaction. He claims that this procedure removes certain interfering substances, notably urochrome.

3. Fermentation Tests.

a. Saccharometer Method. Rub up in a mortar about 15 ml. of urine with a small piece of compressed yeast or with about 0.5g, active dry yeast. Treat in the same way 15 ml. of urine known not to contain glucose and 15 ml. of urine known not to contain glucose and 15 ml. of urine to which glucose has been added. These are necessary controls. Transfer each mixture to a saccharometer (Fig. 15, p. 69) and stand it aside in a warm place for about 12 hours. If glucose is present, alcoholic fermentation will occur and carbon dioxide will coffect as a gas in the upper portion of the tube. On the completion of fermentation, fill the bulb portion of the saccbarometer with 10 per cent sodium hydroxide solution, close the opening of the apparatus tightly with the thumb, mix the contents thoroughly, and restore the gas space to the limb of the saccharometer. Holding the apparatus at arm's length and with face averted (to avoid accidents) remove the thumb from the opening. Remembering that sodium hydroxide has the power to absorb carbon dioxide, how do you explain the result?

Ordinary bakers' yeast (Saccharomyces cerevisiae) will readily ferment glucose, fructose, mannose, maltose, and sucrose. Galactose, lactose, and the various pentoses are not fermentable, or, if so, at such a slow

² Hammarsten suggests that the solution be holled for two to five minutes (according to the sugar content) over a free fiame and the tube then permitted to stand five minutes before drawing conclusions.

^a A dr. powder containing the ingredients for a Nylander test is available commercially under the name of "Galatest" powder (obtamable from the Denver Chenical Mfg Co., Inc., New York City.). In use, a drop of unne is placed on a small amount of the powder. If reducing sugar is present, the powder turns gray or black.

rate as to be of no practical significance in connection with the above test, yeast which readily ferments galactose may be obtained, however, by incubation and growth of ordinary baker's yeast on a medium rich in galactose. Thus the yeast fermentation test is of considerable practical value in distinguishing between glucosuria, lactosuria, and pentosuria, either simple or mixed, particularly when exhaustive yeast fermentation is used in connection with reduction tests before and after treatment with yeast, as in Mathews' procedure which follows

- b Matheus' Modification for Distinguishing Fermentable Sugar (Glucose) from Nonfermentable Reducing Substances (Lactose, etc.) Nix 20 ml of urine in a test tube with 2 3 g of active dry yeast, stirring vigorously Place at an angle of about 45° in a beaker of water and keep at 42° C for 50 minutes, mixing occasionally 11 for cent or more of glucose is present, an evolution of gas bubbles will be observed after a few minutes and all glucose will be decomposed (up to 6 per cent) in 50 minutes. Filter and apply Renedlet s test A positive test indicates the possibility of factose, but only if a control test using the same amount of yeast with urine containing added glucose demonstrates that the yeast is capable of removing all glucose under the experimental conditions. Reducing sugar may also be determined quantitatively before and after the fermentation
 - 4 Polariscopic Examination For directions as to the use of the polariscope, see p 69, and for application to the quantitative analysis of urine Chapter 31

PROTEINS

Normal urine contains a trace of protein material but the amount present is so slight as to escape detection by any of the simple tests in general use for the detection of protein urinary constituents. The following are the more important forms of protein material which have been detected in the urine under pathological conditions.

- 1 Serum albumin
- 2 Serum Llobulin
- Proteoses and peptones

 You blood proteins

 Proteins of Bence-Jones and of Hektoen,
 Kretschmer and Welker
- 4 Nucleoprotein
- 5 Fibrin
- 6 Oxyhemoglobin and related compounds
 - 7 Myorlobin

ALBUMIN

Normal urine contains a trace of albumin which is too slight to be detected by the usual procedures 1thuminaria is the condition in which readily detectable amounts of the serum proteins are found in the urine. The proteinum is most commonly due to the presence of scrim albumin, since albumin is the most abundant of these serum proteins, and has the smallest molecular size thus permitting the greatest diffusion through damaged membranes. It is not uncommon however to find significant amounts of globulin material in so-called albuminum at the proper tests.

are applied. There is some uncertainty as to whether or not the scrum proteins are altered from their normal state when found in the grine

There are two distinct forms of albuminum, viz, renal albuminuma and accidental albuminuma. Sometimes the terms true albuminuma and false albuminuma are substituted for those just given. In the renal type the albumin is exereted by the lidneys. This indicates a more serious condition and at the same time is more frequently encountered than the accidental type. Associated with renal albuminum are usually altered blood pressure or kidney structure. In the accidental form of albuminuma the albumin is not exerted by the kidneys (as is the case in the renal form of the disorder) but arises from the blood, lymph, or some albuminecontaining evidate coming into contact with the urine at some point below the kidneys.

The determination of albumin may be of assistance in following the course of kidney disturbances, but the results can only be interpreted in the light of other chinical findings. Even in nephritis not every sample of urine will be abnormal if the specimens are obtained under ordinary conditions. It is nevertheless rare in this condition to fail to find albumin, casts, or excessive ted cells and white cells in a concentrated urine. No diagnosis of the presence or absence of nephritis should be made until such as

findings are observed

Benign proteinuria, which is usually transitory and in which there is no evidence of permanent ladney damage, may be due to such causes as severe evercise or cold haths. It is particularly common in young people. In the unusual condition known as orthostatic albuminuria, the proteinuria is apparently due to posture, urne formed while the patient is lying down is free from protein, while that formed when the upright position is assumed, particularly the position of military attention, contains protein. This condition is apparently harmless, although it has been used as a deterrent to military service, and may be due, in some instances at least, to mechanical pressure on the renal blood vessels due to a lumbar lordesis.

Foreign proteins injected into the blood stream are excreted by way of the urine, and lead to some excretion of blood proteins also Pathological albuminum may in some cases be due to the passage of an abnormal protein from the tissues into the blood

TESTS FOR ALBUMIN

The urms should be filtered or centrifuged before these tests are performed, and care should be taken against misinterpretation of the presence of pus or bacteria. None of the tests described here is specific for albumin, since practically all of the proteins which have been found in urms at one time or another will respond to most of these tests. It is common clinical practice, however, to refer to 2 positive reaction as signifying the presence of albumin in the absence of specific information that some other protein (globulin, hemoglobin, Bence-Jones protein etc.) is present, since in the vast majority of cases it appears to be really albumin which is concerned A positive test for albumin would entail its characterization is a literacquallable protein, soluble in salt-free water (i.e., after dialysis), and re-

quiring more than half saturated ammonium sulfate for precipitation (see Chapter 6 for further discussion)

The official test for albumin adopted by the Association of Life Insurance Medical Directors of America may be found in Chapter 31

Nitric Acid Ring Test (Heller). Place 5 ml. of concentrated HNO, in a
test tube, incline the tube, and by means of a pipet allow the urine to flow
slowly down the side The liquids should stratify with the formation of a
white zone of precipitated protein at the zone of contact.

If the albumin is present in very small amount the white zone may not form until the tube has been allowed to stand for several minutes. If the urine is quite concentrated, a white zone, due to urie acid or urates, will form upon treatment with nitne acid as indicated. This ring may be easily differentiated from the albumin ring by repeating the test after diluting the urine with 3 or 4 volumes of water, whereupon the ring, if due to uric acid or urates, will not appear It is ordinarily possible to differentiate between the albumin ring and the une acid ring without diluting the urine, since the ring, when due to urie acid, has ordinarily a less sharply defined upper border, is generally broader than the albumin ring, and frequently is situated in the urine above the point of contact with the nitric acid. Concentrated urines also occasionally exhibit the formation, at the point of contact, of a crystalline ring with very sharply defined horders. This is urea nitrate and is easily distinguished from the fluffy ring of albumin If there is any difficulty in differentiation, a simple dilution of the urine with water, as above described, will remove the difficulty Various colored zones, due either to the presence of indican, hile pigments, or to the oxidation of other organic urinary constituents, may form in this test under certain conditions. These colored rings should never he confounded with the white ring which alone denotes the presence of albumin

After the administration of certain drugs, a white precipitate of resin acids may form at the point of contact of the two fluids and may cause the observer to draw wrong conclusions. This ring, if composed of resin acids, will dissolve in alcohol, whereas the albumin ring will not dissolve in this solvent.

A ring closely resembling the albumin ring is often obtained in urines preserved for a considerable time by thymol when subjected to the mirro and test. The ring is due to the formation of mitrosothymol and possibly mitrothymol. If the thymol is removed from the urine by extraction with petroleum ether? previous to adding nitro and, the ring does not form.

 Nitric Acid and Magnesium Sulfate Ring Test (Roberts) Place 5 ml of Roberts' reagent" in a test tube, incline the tube, and hy means of a pipet allow the urine to flow slowly down the side. The liquids should stratify with the formation of a whate zone of precipitated protein between the layers.

Accomplished readily by gently agitating equal volumes of petroleum ether and the urne under examination for five minutes in a test tube before applying the test. is See Appendix.

This test is a modification of Heller's ring test and is rather more satisfactory than that test, since the colored rings never form and consequent confusion is avoided.

- 3. Sulfosolicylic Acid Test. To 1 volume of urine add 2 to 3 volumes of 3 per cent sulfosalicylic acid solution. A turbidity (compare against a control diluted with water) or precipitate denotes the presence of albumin or globulin (but not proteose). The precipitate may be intensified by warming.
- 4. Coogulation or Boiling Test. a. Heat 5 ml. of urine to boiling in a test tube. (If the urine is not clear it should he filtered.) A precipitate forming at this point is due either to alhumin (or globulin) or to phosphates. Acidify the urine slightly by the addition of 3 to 5 drops of very dilute acetic acid, adding the acid drop hy drop to the hot solution. If the precipitate is due to phosphates it will disappear under these conditions, whereas if it is due to protein it will not only fail to disappear but will become more flocculent in character, since the reaction of a fluid must he acid to secure the complete precipitation of the protein by this coagulation process.

Too much acid should be avoided since it will cause the protein to go into solution. Certain resin acids may be precipitated by the acid, but the precipitate due to this cause may be easily differentiated from the protein precipitate by reason of its solubility in alcohol.

- b. A modification of this test in quite general use is as follows: Fill a test tube two-thirds full of urine and gently heat the upper half of the fiuld to boiling, heling careful that this full does not mix with the lower half, which serves as a control. A turbidity indicates protein or phosphates. Acidify the urine slightly by the addition of 3 to 5 drops of dilute acetic acid, whereupon the turbidity, if due to phosphates, will disappear.
- 5. Osgood-Hoskins Test for Urinary Protein. To 5 volumes of urine add 1 volume of 50 per cent acetic acid, followed by 3 volumes of saturated (30 per cent) sodium chloride. (The appearance of a precipitate after the addition of acetic acid, at room temperature, indicates the presence of bile saits, urates, resin acids, etc., whereas a precipitate after the addition of the salt solution suggests Bence-Jones protein (see p. 832), or globulin in excess of 0.38 g. per liter.) Heat the mixture gradually to boiling. As the temperature is raised the precipitate of Bence-Jones protein, if present, will go into solution; if albumin or globulin are present a precipitate will form. This test has the advantage of indicating the presence of Bence-Jones protein as well as albumin and globulin.

GLOBULIN

Serum globulin is not a constituent of normal urine but frequently occurs in the urine under pathological conditions and is ordinarily associated with serum albumin. In albuminurin globulin in varying amounts often accompanies the albumin, and the clinical significance of the two is very similar. Under certain conditions globulin may occur in the urine unaccompanied by albumin.

TESTS FOR GLOBULIN

Globulin will respond to all the tests outlined above under "Albumin." If it is desirable to differentiate between albumin and globulin in any urine, the following procedure may be employed:

Place 25 ml of neutral urine in a smalf beaker and add an equal volume of a saturated solution of ammonium sulfate Globulin it present will be precipitated if no precipitate forms add ammonium sulfate in substance to the point of saturation if albumin is present it will be precipitated upon saturation of the solution as just indicated. This method may also be used to separate globulin and albumin when they occur in the same urine

Frequently in urine which contains a large amount of urates a precipit rate of ammonium urate may occur when it e ammonium sulfate solution is added to the urine. This urate precipitate should not be confounded with the precipitate due to globulin. The two precipitates may be differentiated by means of the fact that the urate precipitate ordinarily appears only after the lapse of several minutes whereas the globulin generally precipitates at once

NONBLOOD PROTEINS

Certain proteins are occasionally exercted by the kidneys which do not give the precipitin reactions for any of the normal blood proteins. These



Fig 227 CRYSTALLINE BENCE-JONES I ROTEIN
Isolated by Summerson from the unne of a case of mul
t ple my cloma

include proteoses Benee-Jones protein and the protein of Hektoen Kretischmer and Welker

Proteoses or substances Living similar precipitation reactions have frequently been found in the urine in cases of picumonia diphtheria intestinal ulcer careinoma dermatitis ostcomalacia atrophy of the kidneys and in conditions in which there is absorption of partially digested pus

Bence-Jones protein is believed to be of diagnostic importance in cases of multiple myeloma and myelogenic osteo-arcoma. It has been shown to be channeally and immunologically distinct from any of the blood proteins and has been obtained in crystalline form (see Fig. 227). It appears

to be present normally in small amount in the hone marrow and in certain white hlood cells. Its exerction in relatively large amount (30 to 50 g. per day in some instances) under pathological conditions is apparently due either to overproduction or decreased utilization; it is not known which is the true explanation. The presence of Bence-Jones protein in the blood plasma of patients with multiple myeloma has been shown by electrophoretic methods

The protein of Hektoen, Kretsehmer, and Welker resembles the Bence-Jones protein and certain proteoses in solubility and precipitation limits with ammonium sulfate (40 to 60 per cent). It is distinct from these, and from blood proteins in its precipitin reactions, its hehavior with heat, and in crystalline form.

TEST FOR NONRLOOD PROTEINS

I. Schulte's Method. Acidify 50 ml, of urine with dilute acetic acid and filter off any precipitate of nucleoprotein which may form. Now test a few ml, of the urine for coagulable protein, by test 4 under "Albumin," p. 831. if coagulable protein is present, remove it by coagulation and filtration before proceeding. Introduce 25 ml. of the urine, freed from coagulable protein, into 150 ml. of absolute alcohol and allow it to stand for 12 to 24 hours. Decant the supernatant fluid and dissolve the precipitate in a small amount of hot water. Now filter this solution, and after testing again for nucleoprotein with very dilute acetic acid, try the bluret test. If this test is positive the presence of proteose is indicated."

Urobliin does not ordinarily interfere with this test since it is almost entirely dissolved by the absolute alcohol when the proteose is precipitated.

- Boiling Test. Make the ordinary coagulation test according to the directions given under "Albumin," p. 831. If no coagulable protein is found, allow the boiled urine to stand and note the gradual appearance, in the cooled fluid, of a flaky precipitate of proteose.
- J. Detection of Bence-Jones Protein. Heat the suspected urine very gently, carefully noting the temperature. At as low a temperature as 40° C. a turbility may be observed, and as the temperature is raised to about 60° C. a flocculent precipitate forms and clings to the sides of the test tube. If the urine is now acidified very slightly with acetic acid and the temperature further raised to 100° C., the precipitate at least partly disappears. Filter while boiling hot. The precipitate returns on cooling the tube. The addition of a few drops of dilute (1 per cent) calcium chioride solution to the urine before testing frequently improves the response to this test.

This property of precipitating at so low a temperature and of dissolving at a higher temperature is typical of Bence-Jones protein and may be used to differentiate it from all other forms of protein material occurring in the Utline.

4. Osgood-Huskins Test for Bence-Junes Prutein. See Exp. 5. p. 831.

If it is considered descrable to test for peptone the proteose may be removed by saturation with (NHa) SO₂ according to the directions given on p 197 and the filtrate tested for peptone by the buyer test.

NUCLEOPROTEIN

There has been considerable controversy as to the proper classification for the protein material which forms the nubecula of normal urine By different investigators it has been called mucin, mucoid, phosphoprotein, nucleoalbumin and nucleoprotein Of course, these terms are not synonymous. Mucin and mucoid are glycoproteins and hence contain no phosphorus (see p. 185), whereas phosphoproteins and nucleoproteins are phosphoproted compounds. It may possibly be that both these forms of protein 1c the glycoprotein and the phosphorized type, occur in the urine under certain conditions (see p. 810). In this connection we will use the term nucleoprotein. The pathological conditions under which the content of nucleoprotein is increased include all affections of the urinary passages and in particular pychitis, nephritis, and inflammation of the urinary bladder.

TESTS FOR NUCLEOPROTEIN

- 1 Detection of Aucleoprotein Place 10 ml of urine in a small beaker, dilute it with three volumes of water to prevent precipitation of urates, and make the reaction very strongly acid with acetic acid if the urine becomes turplid it is an indication that nucleoprotein is present
- If the urine under examination contains albumin the greater portion of this substance should be removed by boiling the urine before testing it for the presence of nucleoprotein
- 2 Tannic Acid Precipitation Test (Ott) Mix 25 mi of the urine with an equal volume of a saturated solution of sodium chloride and slowly add Almen a reagent 11 in the presence of nucleoprotein a voluminous precipitate forms

BLOOD

The patbological conditions in which blood occurs in the urine may be classified under the two divisions hematuria and hemoglobinuma. In hematuria we are able to detect not only the hemoglobin hut the unruptured corpuscles as well whereas in hemoglobinuma the pigment alone is present. Hematuria is brought about through blood passing into the urine because of some lession of the kidney or of the urinary tract below the kidney. Hemoglobinuma is brought about through hemolysis, i.e., the rupturing of the stroma of the erythrocyte and the liberation of the bemoglobin. This may occur in malaria typhoid yellow fever, hemolytic jaundice, and other diseases. It may also occur as the result of a burn covering a considerable area of the body through the action of certain hemolytic poisons or as a result of transfusion with incompatible blood.

DETECTION OF BLOOD

1 Benildine Reaction This is one of the most delicate of the reactions for the detection of blood Different benzidine preparations vary greatly in their sensitiveness however Inasmuch as benzidine solutions change readily upon contact with light it is essential that they be kept in a dark place.

¹ See Appendix.

The test is performed as follows. To 3 ml of a saturated solution of benzidine in glacial acetic acid 12 add 2 ml. of urine and 1 ml. of 3 per cent hydrogen peroxide. A blue or green color indicates a positive test. The following test is more delicate and specific.

Confirmatory Test. To 10 ml. of urine add 1 to 2 drops of glacial acetic acid and extract by shaking with 5 ml. of ether Pour the ether extract into a small evaporating dish. Put on a hot water bath (with fame turned out) and evaporate to dryness. To the residue add a few drops of water, a drop of benzidine solution, and a drop of hydrogen peroxide. A blue or green color indicates blood.

Often when urmes containing a small amount of blood are tested by the direct procedure, the mixture is rendered so turbid as to make it difficult to decide as to the presence of a faint green color. The sensitiveness of the benzidine reaction is greater when applied to aqueous solutions than when applied to the urme.

For modifications of this test and further discussion see page 484

2. Guaiac Test. Place 5 ml. of urine 14 in a test tube and by means of a pipet introduce a freshly prepared alcoholic solution of guaiac (strength about 1 60) into the fluld until a turbidity results, then add old turpentine or hydrogen peroxide, drop by drop, until a hlue color is obtained.

This test is also much more delicate when applied to the acid-ether extract. The test is positive both hefore and after holling the urine for 15 to 20 seconds. Pus does not respond after holling. Old, partly putrefied pus gives the test even without the addition of hydrogen perovide or old turpentine, whereas fresh pus responds upon the addition of hydrogen perovide. See the discussion on p. 480 and the test on p. 484.

3. Spectroscopic Examination. Submit the urine to a spectroscopic examination according to the directions given on p. 494, looking especially for the absorption bands of oxyhemoglobin and methemoglobin (see Fig. 110, p. 473).

MYOGLOBIN

This home pigment, derived from muscular tissue, is found in the urine after extensive destruction of muscular tissue, as from crushing injuries Urine containing myoglobin resembles that containing blood, it may be smoky dark brown or red, and will give a positive benzidine test. Red cells, however, are notably absent, and the sediment may contain brown pigmented easts. Myoglobin has a molecular weight only about one-fourth that of hemoglobin (see Chapter 22 for further discussion) which probably explains its ready diffusibility through the kidney membranes. The appearance of myoglobin in the time is usually associated with concomitant kidney damage, both chinically and experimentally, but whether this has any direct connection with the presence of myoglobin is not known.

¹³ Glacial acetic acid is preferable but if it is not available, alcohol acidified with acetic acid may be used.

¹⁶ Mkaline urme should be made slightly send with acetic acid as the blue end reaction is very sensitive to alkali

PUS

Pus may be present in the urine in inflammatory affections of various types Such a condition is termed pjuria. Ilbumin always accompanies the pus. In catarrh of the urinary bladder and in inflammation of the urithra or of the pelvis of the kidney, pus is particularly apt to be present in the urine. If a urine of high pus concentration is voided it may indicate the rupturing of an abscess in some part of the genitourinary tract. Pus may be detected by one of the procedures given below.

TESTS FOR PUS

I Microscopical Detection of Pus The characteristic form elements of pus are leukocytes. They may occur in very small number in normal urine. Examine the unne (centriloged if necessary) under the microscope Any considerable number of pus corpuscles indicates a patl ological unne. In and urine the pus corpuscles appear as round colories cells composed of refractive, granular protoplasm. Sometimes they may exhibit amchoid movements particularly if the slide containing them be warmed slightly. They are nucleated (one or more nuclei) the nuclei being clearly visible only upon treating the cells with water, acete and or some other suitable reagent in alkaline urine the pus corpuscles are often degenerated. They may occur as wollen, transparent cells withe their it no granular structure. If the degeneration has proceeded far enough the nuclei fade and the cell disintegrates and only debus remains.

Sometimes it is almost impossible to differentiate between pus corpuscles and certain types of epithelial cells. In such a case apply the following chemical test

2 Guazac Test This test is not specific for pus, but is given by certain other substances and particularly by blood (see p 835). Perform the test as follows Acidify the urnse (if alkaline) with acetic acid, filter and add uncture of guazac to the sediment on the paper. If the pus is old and partly purefied, it will give a blue color I in no blue color is secured, add old turpentine, or bydrogen percuide, drop by drop A blue color formed only under these conditions indicates fresh pus.

As a control test boll some of the urine (or sediment) for 15 to 20 seconds and repeat the test. Pus does not respond after boiling in the case of blood the test is positive both before and after boiling.

BILE

Both the pigments and the acids of the bile may be detected in the urine under certain pathological conditions. A urine containing bile may be yell lowesh green to brown in color and when shaken foams readily. The staining of the various tissues of the body through the absorption of bile due to occlusion of the hile duct is a prominent symptom of the condition known as interus or jaundice. Bile is always present in the urine under such conditions unless the amount of bile reaching the tissues is extremely small.

TESTS FOR BILE PIGMENTS

Practically all of the tests for bile pigments are based on the oxidation of the pigment by a variety of reagents with the formation of a series of colored derivatives. For detailed chemistry of these tests see the chapter on bile. A simple though satisfactory test for bile consists in shaking the uries in a test tube and observing the yellow foam.

- 1. Gmelin's Test. To about 5 ml. of concentrated nitric acid in a test tube add an equal volume of urine carefully so that the two fluids do not mix. At the point of contact note the various colored rings; green, blue, violet, red, and reddish yellow. (Fuming yellow nitric acid gives the best results.)
- Rosenboch's Modification of Gractin's Test. Filter 5 ml. of urine through a small filter paper. Introduce a drop of concentrated nitric acld into the cone of the paper and observe the succession of colors as given in Cmelin's test.
- 3. Harrison Spot Test. 12 & S filter paper No. 470 is impregnated with a 10 per cent barium chloride solution and dried. Dip the dry paper into the urine for 10 seconds and add a drop of Fouchet's reagent" at the surface line. A green color indicates billrubin. This test bas been modified by Hawkinson, Watson and Turner. 12

TESTS FOR BILE ACIDS

- I. Furfural-H₂SO, Test (Mylius). To 5 ml, of urine in a test tuhe add 3 drops of a very dilute (1:1000) aqueous solution of furfural. Now incline the tube, run ahout 2 to 3 ml. of concentrated suifuric acid carefully down the side, and note the red ring at the point of contact. Upon slightly agitating the contents of the tube the whole solution gradually assumes a reddish color. As the tube becomes warm, it should he cooled in running water in order that the temperature may not rise above 70° C.
- It is claimed that this test is not satisfactory in the presence of protein and chromogenic substances which yield interfering colors with sulfuric acid.
- 2. Surface Tension Test (Hoy). This test is hased upon the principle that bile acids have the property of reducing the surface tension of fluids in which they are contained. The test is performed as folions: Cool about 10 ml. of fresh urine in a test tube to 17° C. or lower, and sprinkle a little finely pulverized sulfur upon the surface of the fluid. The presence of bile acids is indicated if the sulfur sinks to the bottom of the liquid, through the surface film. Compare with a control tube of normal urine known to be free from bile. Urines preserved with thymol may respond positively to this test.

THE ACETONE BODIES

The accione (or ketone) bodies include the compounds acctoacetic (or diacetic) acid, β -hydroxybutyrie acid, and acctone. The chemical relationship between these various acctone bodies is as indicated below; acctoacetic acid and β -hydroxybutyric acid are primary products, the latter probably being formed by reduction of the former. Acctone, however, is a decomposition product of acctoacetic acid and is probably not produced as such within the body, although it is invariably found there when the other two acctone bodies are present.

CII₂ CO CII₂ COOH → CH₂ CO CII₂ + CO
Acetoacetic acid Acetone

| II (reduction)

CH₂ CHOΠ CH₂ COOH
β-Hydroxybutyric acid

nonvolatile compound with bisulfites

rent of air under the proper conditions

Acetoacctic acid and β-hydroxybutyrie acid appear to be either intermediate products in the breakdown of fatty acid chains or secondary combinations of 2-carbon fragments formed in this breakdown or closely related to it Under normal conditions the fatty acids in the animal body are oxidized completely to carbon dioxide and water and intermediate products do not appear to any great extent in the blood or urine In certain abnormal conditions, however, the ketone bodies accumulate in the blood (letonemia) and are exercted in the urine (letonuria), this general condition is known as a kelosis Ketosis is apparently always associated with some abnormality of earbohydrate metabolism. It is still a disputed question whether ketosis results from the failure on the part of the animal body to oxidize completely the ketone hodics produced normally, or from an overproduction of ketone bodies by the organism in an attempt to meet fuel requirements not supplied by carbohydrates. Ketosis occurs in fasting or during carbohydrate deprivation and a such conditions disappears when carbohydrate is fed Pathologically it is most severe in diabetes mellitus, in which disease the accumulation of the ketone body acids is largely responsible for the development of diabetic acidosis Diahetic urine during ketonuria is often high in ammonia which is formed by the body to combat acidosis Diabetic ketosis is alleviated by insulin treatment Letosis also occurs in the acetonemic vomiting of childhood, and frequently in pregnancy, fevers, ether and chloroform anesthesia, mal nutrition, prolonged feeding of a earbohydrate-poor diet high in meat and fat, and many other conditions (See also Chapters 31 and 33)

Acetone, CH₂ CO CH₂. Acetone when pure is a liquid which possesses a characteristic aromatic fruitlike odor, boils at 50° to 57° C, and is muscle be with water, alcohol, or ether in all proportions it reacts with alkhland iodine to yield iodoform. With sodium introprusside in alkaline solution it gives a red ecompound. With saleylie aldchyde, in strongly alkaline solution, it forms a red to orange condensation commound it forms.

Acetoacetic Ácid, CII, CO CII, COOII. In the pure state acetoacetic acid is a colorless liquid miscible with water, alcohol, and ether, in all proportions. Its solution gives a Bordeaux-red color with ferric chloride (not given by acetone). With introprusside in alkaline solution it gives a permanganate color, similar to that given by acetone but many times more intense on an equivalent basis. In aqueous solution acetoacetic acid decomposes (most readily on warming in acid solution) to yield acetone. Hence after heating or long standing it gives the tests for acetone, and is probably never found in urine without the concomitant presence of actione. It is always determined quantitatively as acetone. Acetone and

acctoacctie acid may be separated by blowing off the acctone with a cur-

 β -Hydroxybutyric Acid, CH₂ CHOH CH₂ COOH. Ordinary β hydroxybutyric acid is an odorless, transparent, levorotatory syrup easily soluble in water, alcohol, and ether, it may be obtained in crystalline form it does not respond to the tests for acetone or acetoacetic acid unless it is first oudized, when it yields acetoacetic acid and, on decomposition, acetone β -Hydroxybutyric acid in acid urines acts as a urinary antiseptic Hence ketogenic diets may be useful in cases of infection of the urinary tract

Occurrence of Acetone Bodies in Urine. Total ketone bodies are found in normal urine to the extent of about 20 mg (expressed as acetone) in 24 hours, but this amount is variable Of the total, one-balf or more is usually in the form of β hydroxybutyric acid, but this ratio is also variable Patbologically, values of from 0 02 to 6 g or more per day of combined acetone and acetoacetic acid have been observed β -Hydroxybutyric acid has been found in the urine in severe diabetes in amounts of 50 to 100 g or over In such conditions the β hydroxybutyric acid may constitute 60 to 80 per cent of the total acetone bodies. In rare cases excretion of large amounts of β -hydroxybutyric acid may occur with low acetone output

TESTS FOR ACETONE BODIES

The three acetone bodies practically always occur together in the urms and bare essentially the same significance. The usual tests for letonuria are tests for acetone or acetoacetic acid or both, since a distillate obtained by beating urine will contain acetone derived by decomposition of acetoacetic acid as well as preformed acetone, and the introprusside reaction applied to urine will detect either compound, although with quite different relative sensitivity. There is no satisfactory simple direct test for β hydroxybutyric acid in urine. The ferric chloride test for acetoacetic acid is not given by acetone, but the test is neither sensitive nor specific and caunot be recommended for chiucal work, except perhaps as a confirmatory test. It must be made on the urine directly, and not on a distillate

The introprusside test (particularly the Rothera version) is much more sensitive (at least five or more times) to acetoacetic acid than to acctone, hence when applied to urine directly the result will depend largely on the relative amounts of these two substances present. Fresh urine may give a positive test, the same urine on standing and after decomposition of the acetoacetic acid to acctone may give a negative test. In a distillate, where both compounds are present as acctone, the reaction is sometimes fainter than in the original urine. The test is not entirely specific for the acctone bodies, but when positive it is strongly indicative of their presence. It is probably the most widely used test for ketonuria in chinical work, and many variations in technique have been proposed, nevertheless it cannot be regarded as entirely satisfactory.

The iodoform and salicylic aldehyde tests react with acetone but not directly with acetoacetic acid, and because of interfering substances in urine must be carried out on acetone separated from the urine by distillation or otherwise. The iodoform test on a distillate is quite sensitive and may be of value in suggesting the presence of very small amounts.

of acctone bodies, but it is not entirely specific for acctone (alcohol for cample also gives a positive test) and it is less satisfactory for clinical work than the other tests described here. The salicylic aldehyde test, carried out as described here on acctone separated from the urine, is probably the most generally satisfactory test for ketonuma.

TESTS FOR ACETONE AND ACETOACETIC ACID

- 1. Isolation from the Utine The teste for acetone are more satisfactory if the acetone is separated from the urine by distillation and the tests applied to the distillate Introduce into a small distilling built 10 ml (or more) of the urine to he tested, and acidify Distil off about 2 ml. of liquid (containing most of the acetone) into a test tube A condenser is not necessary but the tube should be kept cool. In the distillation any acetoacetic add present is decomposed and the acetone from this source also passes into the distillate Try either of the two following tests on the distillate.
- 2 Iodoform Test (Lieben) To 2 ml of distillate add 3 to 5 drops of 10 per cent NaOll and then Lugol s todne solution drop by drop to a faint yellow Let stand at room temperature if necessary A definite turbidity changing to a yellow precipitate of lodoform sbould be noted. The odor is characteristic as is also the crystalline form viewed under the microscope (see p. 68).

This test is given by alcohol which may be formed by fermentation in diabetic urlines. Alcohol, bowever, reacts much more slowly. If ammonia is used, usually 5 to 10 drops, instead of NaOH (Gunning's test) the reaction is more specific for acetone but is less delicate.

- 3 Nitroprusside Test (Legal) Try this test on both the original urine and the distillate To 2 ml of liquid add a few drops of a freshly prepared 5 per cent aqueous solution of sodium nitroprusside Make alkaline with Aolli A red color Indicates acetone If the test is made directly on urine a red color is given by creatinine which, however, disappears on the addition of acetic acid A modification of this test in quite general use is as follows to a few ml of urine add a few drops of the introprusside solution and mix. Add concentrated ammonium hydrounde carefully down the sade of the tube so as to form a layer over the sample. A purple (not hrown) ring at the zone of contact indicates the presence of acetone bodies.
 - 4 Miropruside Test (Rothers). Saturate 20 ml of urine with ammonium sulfate by shaking with the crystals in a test tube. Add 2 to 3 drops of concentrated Nil,Oll and a few drops of a freshly prepared 5 per cent solution of sodium nitroprusside and shake. A positive test is indicated by the development of a permanganate tinge which gradually deepens. A brown color is not a positive test. A quick strong reaction indicates shout 0.25 per cent acetoacetic acid while a slow weak reaction is given by 0.005 per cent acetoacetic acid. A faint test has less significance than the ferric chorde test? because of the delicacy of the reaction. The test is given by acetone also but is much less delicate for this substance.
 - 5 Salicylic Aldehyde Test (Behse) = In this test a previous distillation is unnecessary since the acetone, preformed and from acetoacetic acid, is dis-

u Tie dry ingredients of the Roll era test for acctone are available commercially as a powder (Arctione Test | Itamable from the Denver Chemical Mig Co Inc \ver York City) in use a few dry so during an epiaced on a small amount of the powder A purple color indicates acctring.

Behre J Lab Clin. Med. 12, 770 (1928) Also personal communication.

tilled in the test tube onto the reagents. Place 3 mi. of uriue in a clean test tube and 3 ml. of distilled water as a contral in another tube, and add 1 drop of 1:1 sulfuric acid to each, Prepare two small thin squares of cotton and in the center of each place a drop of salicylic aldehyde (or of an alcoholic solution of salicylic aldehyde)) and two drops of a saturated solution of potassium hydroxide. These reagents solidlify to form a yellow dlsk. When they have solidlified, invert a cotton square over each test tube and push in slightly so that the spot of reagents faces dnwn tuward the fiuld in the tube but does not touch the sides of the tube. Place both tubes upright in boiling water for eight minutes. Remove the cotton and examine the spots. The presence of acetone or acetoacetic acid in the urine is indicated by a plnk to deep rose coloration of the spot, as compared with the yellow color of the blank test. The color from acetone bodies deepens on standing and in doubtful cases should be examined after a few minutes. Familiarity with the test makes it possible to estimate roughly the amount of total acetone present.

6. Ferric Chloride Test for Acetoocetic Acid¹¹ (Gerhordt). To 5 ml. of urine in a test tube add ferric chioride solution, drop by drop, until no more precipitate forms. In the presence of acetoacetic acid, a Bordeaux-red color is produced; this color may be somewhat masked by the precipitate of ferric pbosphate, in which case the fluid should be filtered.

A positive result from the above manipulation simply indicates the possible presence of acetoacetic acid. Before making a final decision regarding the presence of this substance make the two following control experiments:

- a. Place 5 ml. of urine in a test tube, small beaker, or Erlenmeyer flask and boil it vigorously for three to five minutes. Cool the vessel and, with the boiled urine, make the test as given above. Compare with the test on the unboiled sample. As has been already stated, acetoacetic acid yields acetone upon decomposition and acetone does not give a Bordeaux-red color with ferric chioride. By boiling as indicated above, therefore, any acetoacetic acid present would be decomposed into acetone and carbon dioxide and the test upon the resulting fluid would be negative. If positive, the color is due to the presence of substances other than acetoacetic acid.
- b. Place 5 ml. of urine iu a test tube, acidify with H-SO_i, to free acetoacetic acid from its salts, and carefully extract the mixture with ether by shaking. If acetoacetic acid is present, it will be extracted by the ether. Now remove the ethereal solution, evaporate it the dryness, dissolve the residue in 1 to 2 ml. of water, and add 3 to 5 drops of 3 per cent ferric chloride. Acetoacetic acid is indicated by the production of the characteristic Bordeaux-red color.

This color disappears spontaneously in 2½ to 48 hours. Such substances as antipyrine, acetophenetidine, salieyite acid, salicylates, sodium acetate, thocyanates, and thalin yield a similar red color under these conditions; when due to the presence of any of these substances, however, the color does not disappear spontaneously but may remain for days. Many of these disturbing substances are soluble in benzene or chloroform and may

¹¹ Eastman's technical grade or Limer and Amend's And Salrejious, Synthetic, are usually satisfactory, for use without thiston. If Backening appears on the cotton during the heating a solution of 1 part salrejic aldehyde in 1 part methyl or ethyl alcohol should be used.

¹¹ To prepare a solution which may be added to urine, if urines containing this acid is not no tailable for student work, proceed as follows. Treat 12 g, of etb)1 acctoactate with 500 ml. of 0.2 N solumi hydroxide. Allow to stand for 48 hours to hydrolyze the exter In preparing urine for tests and 1 part of this solution to 10 parts of urine.

be removed from the urine by this means before extracting with ether as above. Acutoacetic acid is insoluble in benzene or chloroform

TESTS FOR \$\beta\$-HYDROXYBUTYRIC ACID

1. Black's Reaction. Inasmuch as the urinary pigments as well as any contained sugar or acetoacetic acid will interfere with the delicacy of this test when applied to the urine directly, the following preliminary procedure is necessary Concentrate 10 ml. of the urine under examination to one-third or one-fourth of its original volume in an evaporating dish at a gentle heat Acidify the residue with a few drops of concentrated hydrochloric acid, add sufficient plaster of Paris to make a thick paste, and allow the mixture to stand until it begins to set. It should now be stirred and broken up in the dish by means of a stirring rod with a blunt end. Extract the porous meal thus produced twice with ether by stirring and decantation. Any 8-hydroxybutyric acid present will be extracted by the ether. Evaporate the ether extract spontaneously or on a water bath, dissolve the residue in water, and neutralize it with barrum carbonate To 5 to 10 ml of this neutral fluid in a test tube add 2 to 3 drops of ordinary commercial acid hydrogen peroxide. Mix by shaking and add a few drops of Black's reagent,21 Permit the tube to stand and note the gradual development of a rose color which increases to its maximum intensity and then gradually fades.24

In carrying out the test care should be taken to see that the solution is cold and approximately neutral and that a large excess of hydrogen peroxide and Black's reagent are not added. In case but hittle θ hydroxybutyre acid is present the color will fail to appear or will be but transitory if the existing agents are added in too great cross. It is preferable to add a few drops of the resgent and at intervals of a few innutes repeat the process until the color undergoes no further increase in intensity. One part of θ hydroxybutyre acid in 10,000 parts of the solution may be detected by this test.

2. Polariscopic Examination Subject some of the urine (free from protein) to the ordinary fermentation test (see p. 827). This will remove glucose and fructose, which would interfere with the polariscopic test. Now examine the fermented fluid in the polariscope and if it is levorotatory the presence of \$\tilde{\t

GLUCURONIC ACID

Glueurome acid does not occur free in urine, but is found in combinations known as conjugate glucuronates or glucuronides with a wide variety of compounds under both normal and pathological conditions or after administration of such compounds for medicinal or experimental purposes There are two general types of conjugate glucuronates, the glycoside

¹³ Made by dissolving 5 g. of ferrie chloride and 0 4 g. of ferrous chloride in 100 ml of

²⁴ This disappearance of color is due to the further oxidation of the acetoacetic acid

type and the ester type, both of which involve linkage with glucuronic acid through the OH group on carbon atom number 1 of the cyclic structure. The glycoside type involves the OH group of aliphatic or aromatic alcohols, e.g., phenol, naphthol, horneol, etc., while the ester type involves reaction with the COOH group of such compounds as benzoic acid, phenylacetic acid, etc. Free glucuronic acid is as powerful a reducing substance as glucose, and its quantitative determination may be based upon this fact, ester-type glucuronides likewise show direct reducing properties, probably due to concomitant hydrolysis, while the glycoside type is non-reducing until hydrolyzed Conjugate glucuronates may therefore interfere with the reducing tests for sugar in urine under certain conditions, but they may be readily distinguished from glucose because of their nonfermentability with yeast. While glucuronic acid is destrorotatory, the glucuronides as a class are levorotatory, this also serves for distinction from glucose in urine.

The total glucuronuc acid content of normal urine appears to approximate 0.5 to 1.0 g per day, carlier estimates were considerably lower than this, probably hecause the methods were hased on the mistaken assumption that all conjugate glucuronates are soluble in ether. The glucuronides of normal urine include combinations with phenol, indoxyl, shatoxyl, and the estrogeme hormones. Urine glucuronide content may be greatly increased by the administration of a variety of compounds, such as anti-pyrine, acetylsalicylic acid (aspirin), horicol, camphor, chloral hydrate, menthol, morphine phenolphthalein turpentine, and practically all of the sulfonamides (evcept possibly sulfanilamide). Experimentally, certain carcinogenic substances or derivatives are ultimately evcreted in part as glucuronides. Glucuronic acid formation and conjugation appear to tako place in the hyer. The origin of glucuronic acid is obscure, although it is known to be an end product of the polysaccharide-sphtting action of the enzyme hyaluronidase.

TESTS FOR GLUCURONATES

- I hophthoresorcinol Reaction (Tollens) introduce 5 ml of urine in a test tube and add 0.5 to 1 ml of a 1 per cent solution of naphthoresorcinol in 95 per cent alcohol, and 5 ml of concentrated hydrochloric acid Raise the temperature gradually to the boiling point and boil for one minute, shaking the tube continuously Stand the tube saide four minutes, then cool under the tap Extract with an equal volume of ether (preferably peroide free) Glucuronates are indicated by the ether extract assuming a violeted color The spectroscope shows this extract to possess an absorption band in the green to yellow region of the spectrum. The peak absorption is at approximately 570 min.
- 2 Preparation of Clucuronic Acid (Method of Quick) ³¹ Give 5 g of pulserized borneoi daily to each of several dogs. Collect the urine, acidlfy with acetic acid, and add lead acetate. Most of the coloring matter is carried down kitter, heat the filtrate to boiling, and add an excess of zinc acetate. Filter off the precipitate and wash with but water until no more coloring matter is extracted. This is practically pure zinc borneol glucuronic acid. About 1 g is obtained for each g of borneol given.

¹⁶ Quick J Biol Chem 74 331 (1927)

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Dissoive the finely powdered zine sait in hot 3 5 % sulfuric acid, using about 140 mi for each 100 g of the salt When completely dissolved, cool rapidly and put in an icebox for several hours. Fliter off the crystals of horneol giucuronic acid, wash with a little cold water, and dry in the air

Dissolve 100 g of borneoi glucuronic acid in 1,500 mi of 0 2 \ sulfurle acid and boli for three hours beneath a reflux condenser. Filter and treat the hot filtrate with sufficient barium hydroxide to precipitate tile iast traces of sulfuric acid Allow the mixture to settle Siplion off the supernatant fluid, and complete the separation by centelfuging Concentrate the solu tion under diminished pressure to a syrupy consistency, and let it stand to crystallize Filter off the crystals and wash them with a small amount of alcohol to remove the pigment. This is a mixture of glucuronic acid and its factone Treat 4 g of the product with 200 mi of 95 per cent alcohol and set aside for 12 hours Repeat twice, using 100 mi portions of alcohol The residue should be glucuronic acid of 99 per cent purity. To obtain the pure factone, dissoive some of the mixture of acid and factone in hot giacial acetic acid, allow the solution to cool, and recrystailize from hot water

Glucuronic acid is a syrupy liquid, readily soluble in water and slightly soluble in alcohol. When the aqueous solution is boiled evaporated, or even allowed to stand at the ordinary temperature the acid loses the elements of water and yields the anhydride or lactone. It is a strong or cance acid ($l_x > 1 \times 10^{-3}$)

Glucuronic anhydride CellaO4, forms monochnic tables or needles, having a sweet taste, and m p 160° when heat is gradually applied, or at 170° to 180° when heated rapidly The anhydride is insoluble in alcohol, but dissolves readily in water to form a devirorotatory solution, [a]p =

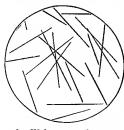


FIG 228 1 ENTOSAZONE CRYSTALS

Isolated and purified in the senior author's laboratory by Dr B L Fleming For color of crystals see Plate II opposite

with an initial value of $[\alpha]_{D}^{*0} = +$ 16° It reduces Fehling's solution on heating, and precipitates the metals from bot alkaline solutions of silver, mercury, and bismuth

PENTOSES

+19 25° The solution prevents the precipitation of cupric ion by alkalies and powerfully reduces Fehling's solution the copperreducing power being 988, com pared with glucose as 100 Glucuronie acid itself is dev trorotatory ($[\alpha]_D = +36^\circ$), but many of its compounds are levorotatory It shows mutarotation

There are two distinct types of pentosuria, viz (1) alimentary pentosuma, resulting from the ingestion of large quantities of

pentose rich fruits such as prunes cherries grapes or plums and fruit juices in which condition the pentoses appear only temporarily in the urine and (2) the chrome form of pentosuria, in which the output of pentoses bears no relation whatever to the quantity and nature of the pentose content of the food eaten Pentosuria is apparently a harmless abnormality; it is definitely known that pentosuria bears no relation to diabetes mellitus and there is no generally accepted theory to account for the occurrence of the chrome form of pentosuria. The pentose detected most frequently in the urine in chrome pentosuria is L-xyloketose. L-Arabinose has been found in cases of alimentary pentosuria. For pentosucione crystals, see Fig. 228.

TESTS FOR PENTOSE

1. Benzidine Reoction (Tauber).²⁶ To 0.1 ml. of the urline in a test tube, add 0.5 ml. of a 4 per cent solution of benzidine in glacial acetic acid. Heat to vigorous boiling, cool under the cold-water tap, and add 1 ml. of water. The presence of pentose is indicated by the immediate appearance of a pink to red color. If pentoses are absent, the mixture has a veilowish-brown color.

This test is highly specific for pentoses in the free form but is not given by gum arabic, nucleic acids, or ribofiavin in which the pentose is combined. Normal and ahnormal constituents of urine do not interfere. Too large amounts of protein may be removed from pathological urine by mixing with an equal volume of 10 per cent trichoroacetic acid solution, warming to 95°, and filtering. The test is applied to the filtrate. This test is said to detect the presence of aldopentoses but not ketopentoses. The 4 per cent benzidine solution is stable for four days.

2. Orcinol-Hydrochloric Acid Reoction (Bial). To 5 ml. of Bial's reagent" in a test tube, add 2 to 3 ml. of urine and heat the mixture gently until the first bubbles rise to the surface." Immediately or upon cooling the solution becomes green and a flocculent precipitate of the same color may form.

This test is believed to be more accurate than the original orcinol test. It is claimed that urines containing menthol, kreosotal, etc., respond to the old orcinol reaction, but not to Bial's. If so desired the osazone of the pentose (see Fig. 228) may be formed, then distilled with bydrochloric acid and the distillate tested by Bial's test (Joiles).

- 3. Phloroglucinol-Hydrochloric Acid Reaction (Tollens). To equal volumes of urline and hydrochloric acid (sp. gr. 1.09) add a little phloroglucinol and heat the mixture on a bolling water bath. Pentose, galactose, or glucuronic acid will be indicated by the appearance of a red color. To differentiate between these compounds examine by the spectroscope and look for the absorption band in the yellow region of the spectrum given by pentoses and glucuronic acid, and then differentiate between the two latter compounds by the melting points of their osagones.
- 4. White-Green Reaction. To the urine in a test tube add two drops of acetic acid and boll. Add Norlt, shake two minutes and filter. To 2 ml. of filtrate add 2 ml. of acetic acid and 5 drops of redistilled anlline. Heat to boiling, allow to stand 2 minutes then cool and extract with 2 ml. of chloro-

¹⁴ Tauber: Proc. Soc. Expd. Biol. Med., 37, 600 (1937).

See Appendix.

¹⁶ The test may also be performed by adding the urine to the hot reagent. No further heating should be necessary if pentose is present.

[&]quot; White and Green: Trans Roy, Soc. Section V. Biological Sciences, Third Series, 24, 145 (1932).

form In the presence of pentose the chloroform extract becomes bright red Glucose and galactose give a green color, fructose a pale yellow, and glucuron ate a very pale yellow

LACTOSE

Lactose is rarely found in the urme except when exercted by women during pregnancy, during the nursing period, or soon after wearing 1c, when the mammary glands are actively functioning. I actosima has no pathological significance representing as it does simply the exerction of a sugar which has found its way into the blood stream by some means and which is not utilizable as such by the organism 10 It is nevertheless important to be able to identify the condition of lactosuria and to distinguish it from glucosuria, otherwise serious misinterpretation of labora tory data may result Lactose in urine is readily distinguished from glucose by its lack of fermentability with ordinary yeast qualitative or better quantitative determination of reducing power before and after exhaustive treatment with yeast is of value in this connection, particularly for the detection of a concomitant glucosuria The characterization of lactose as its typical osazone is not ordinarily attended with much success in urine, better results may be obtained by adsorption of the lactose on charcoal prior to testing as described below. The mucic acid test which is specifie for lactose and galactose may be used to identify lactose in urine but interpretation of results may be obscured by the presence of other insoluble crystalline substances

TESTS FOR LACTOSE

- 1 Fermentation Test Ferment the urine as in Exp 3(a) or 3(h), p 827 If nonfermentable reducing sugar is found it is very probably factose, especially if the patient be pregnant or factisting in rare cases it may be pentose Lac tose may be distinguished from pentose by proper application of Tauber's test (p 845) the mucic acid test or the osazone test.
- 2 Oszone Test (According to Cole) Shake 25 ml of urme with 1,8 of Nierks medicinal charcoal boul few seconds cool thoroughly and shake at intervals for 10 minutes. Filter through a small paper or use a filter pump Let drain completely. Transfer the charcoal to a dish containing 10 ml of water and 1 ml of glacial acettle acid Buil for about 10 seconds and filter hot into a tube containing as much phenylhydrazine hydrochloride as will lee on a quarter and twice as much sodium acetate. Wilk and heat on a bolling water bath for 45 minutes. Remove and let stand for at least an hour. Look for hedgehog crystals of lactoszatone Glucose, if present in significant amount will be adsorbed by the charcoal to some extent and come through along with the lactose to give glucoszone crystals. The value of the charcoal separation is chiefly to increase the concentration of lactose relative to glucose and to facilitate the obtaining of characteristic lactoszone crystals.
 - 3 Mucic Acid Test Transfer 50 ml of urine to a 150 ml beaker and add 12 ml of concentrated th O. Heat on a boiling water bath until the volume is reduced to about 10 ml Cool Add 10 ml of water and let stand overnight A fine white precipitate of mucic acid will form if factose or galactose is

is if any lactose is excreted into the intestinal tract and there by drolyzed into glucose and galactose these products are capable of absorpt on an i utilization by the body

present Examine the crystals under the microscope (see Fig. 19). Other reducing sugars do not give this test. Lactose and galactose may be differentiated by Tollens' test (see tests for pentose)

GALACTOSE

Galactose bas occasionally been detected in the urine, and in particular in that of nursing infants afflicted with a deranged digestive function. It may be present in significant amount in liver disease after the administration of large doses of galactose, as in the galactose tolerance test (analogous to the glucose tolerance test) for liver function. Lactose and galactose may be differentiated from other reducing sugars which may be present in the urine by means of the mucic acid test. For a description of the mucic acid test, see the experiments on lactose above. To differentiate galactose, use Tollens' reaction (see the section on pentose above). The red solution given by galactose shows no absorption bands. Galactose is fermentiable very slowly or not at all by ordinary bakers' yeast.

FRUCTOSE

The occurrence of fructose in the urine is relatively rare. In essential fructosuma small amounts of fructose are constantly exercted regardless of the fructose content of the diet, although it is curious that, if the diet is carbohydrate-free, the exerction of fructose ceases. No satisfactory explanation has as yet been offered for the condition of essential fructosuma, which is apparently a barmless metabolic abnormality. In diabetes mellitus, fructose may at times be excreted along with glucose (never by itself), the significance of this is not known, although it may be recalled that fructose diphosphate is an intermediate in earbobydrate metabolism (see Chapter 33)

TESTS FOR FRUCTOSE

I Borchardt's Reoction. To about 5 ml of urine in a test tube add an equal volume of 25 per cent hydrochloric acid and a few crystals of resorcinol Heat to boiling, and after the production of a red color, cool the tube under running water and transfer to an evaporating dish or beaker Make the mixture slightly alkaline with solid potasslum hydroxide, return it to a test tube, add 2 to 3 ml of ethyl acetate, and shake the tube vigorously in the presence of fructose the ethyl acetate is colored yellow

The only urnary constituents which interfere with the test are nitrites and indican, and these interfere only when they are simultaneously present. Under these conditions, the urnie should be acidified with acetic acid and heated to boiling for one minute to remove the nitrites. In case the indican content is very large, it will impart a blue color to the ethyl acetate, thus masking the yellow color due to fructose. When such urnies are to be examined, the indican should first be removed by Obermayer's test (see p. 804). The chloroform should then be discarded, the acid-urnie mixture diluted with one-third its volume of water, and the test applied as described above. The urnie of patients who have nigosted santonin or rhubrib responds to the test. The test will serve to detect fructose when premet in a dilution of 1. 2000—1 e., 0.05 per cent.

- 2 Resorcinol-Hydrochloric Acid Reaction (Selisanoff). To 5 mi. of Sellvanoff's reagent¹¹ in a test tube add a few drops of the urine under examination and heat the mixture to boiling, or place in a boiling water hath. The presence of fructose is indicated by the production of a red color which ma) or may not lead ultimately to the separation of a red precipitate. The latter if formed may be filtered off and dissolved in alcohol to which it will impart a striking red color.
 - If the boiling be prolonged, a similar reaction may be obtained with urine containing gluco-e The precautions necessary for a positive test for fructose are as follows The concentration of the hydrochloric acid must not be more than 12 per cent, the reaction (red color) and the precipitate must be observed after not more than 20-30 seconds of boiling, glucose must not he present in amounts exceeding 2 per cent, the precipitate must be soluble in alcohol with a bright red color

3. Aminoguanidine Test (Tauber) See p 73

- 4. Phenylhydrazine Test. Make the test according to directions under "Glucose," Exp 1, p 824 With methylphenylhydrazine fructose gives crystals differing from those given by glucose in their rate of formation (p. 73).
 - 5. Polariscopic Examination A simple polariscopic examination, when taken in connection with other ordinary tests, will furnish the requisite data regarding the presence of fructose, provided fructose is not accompanied by other levorotatory substances, such as conjugate glucuronates and \$-hydroxybutyric acid.

ARSENIC

When any soluble form of arsenic is introduced into the body in any way, it is quickly absorbed and distributed by the blood and lympb The absorption is influenced by the quantity and character of food in the stomach, and the activity of the circulation of the part in contact with the poison Some of the absorbed arsenic may be returned to the alimentary canal by way of the bile and gastrointestinal mucous membrane After absorption it may be deposited in the liver, kidneys, brain, bone, muscles, and walls of the stomach and intestines. It is eliminated in all of the excretions, but chiefly by the kidneys and through the feces It does not appear very promptly in the urine but continues to be excreted in the urine over a long period of time, in some cases for several months Testing for arsenic has become of increasing importance because of the widespread use of arsenicals in chemotherapy Many instances of clinical manifestations of arsenical poisoning have been reported

DETECTION AND ESTIMATION OF ARSENIC

1. Gutzett Method " Principle The presence of arsenic is revealed by the brown stain produced on mercuric bromide paper when arsine is liberated from a test solution by the action of nascent hydrogen

²¹ See Appendix

³⁵ Throughout this determination care must be taken to use clean amenic-free reagents

Procedure. Prepare a generator as shown in Fig. 229 using a 50-ml. widemouth hottle fitted with a perforated rubher (As-free) stopper. In the central

chamber (about i cm. wide and 7 cm. long) place loosely packed glass wool or cotton moistened with 10 per cent lead acetate solution. The crit tube should have an internal diameter of about 3 mm. Into this tube insert

a strip of mercuric bromide test paper.

To prepare the arsine stain place 5 ml. of 15 per cent potassium lodide, 5 ml. of acid stannous chloride solution (1.6 g. SnCh dissolved in 100 ml. 10 per cent HCl), 2 ml. of the test solution, and 30 ml. of water in a flask. Add 1.5 g. granulated As-free zinc and stopper immediately with the exit tuhe arrangement. Immerse the generator hottle in water at 25° C. for 1 hour. Remove the test strip and compare the color and length of stain with strips similarly prepared from standard arsenic solutions.

Depending upon the nature of the material under examination the test solutions may be prepared by direct solution or by digestion (tissues, foods, urine) with As-free concentrated sulfuric acid as in the Kjeldahl nitrogen determination (see p. 874). To prepare the standard orsenic solutions dissolve 100 mg. arsenic trioxide powder in 5 ml. of 20 per cent NaOH, neutrolize with dilute HiSO, add 10 ml. excess acid ond make up to 1000 ml. Prepare dilutions of this stock solution with dilute HiSO, so that 2 ml. contain 1, 2, 3, 4 etc. pg. AsiO. These amounts will yield a series of stains of graduated length and intensity.

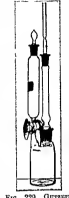


FIG 229 GUTZEIT ARSENIC APPARATUS Courtesy, 4 H. Thomas Company, Philadelphia

2. Reinsch's Test. This test is very much simpler, but not so delicate. It has the advantage of application in the presence of organic matter. The test is performed as follows: The urine, acldified with one-fifth its volume of pure hydrochloric acid, is piaced in a heaker. A piece of bright copper foil free from arsenic is then introduced, and the urine is heated almost to the boiling point. It is then set aside for six to eight hours. The arsenic is deposited on the copper foil, giving it a hluish-gray color. The foil is then removed, washed successively in pure water, alcohol, ether, and dried without heat. The foil is then rolled into a scroll and inserted into a 3-mm.-hore glass tube 4 inches long, about 1 inch from the end. The tube is then held in the Bunsen flame at an angle of 20 to 25 degrees, heat heing applied where the copper foil is situated. The arsente volatilizes and is oxidized, and deposits as octahedral crystals of arsenic trioxide on the cooler part of the tube. The crystals can readily he recognized by the microscope and sometimes with a simple magnifying lens.

and glassware A blank determination should be run on the reagents in each generator used in a series. For the precise quantitative determination of arsenic by the Gutzeit method consult Victhods of Analysis of the Association of Official Agricultural Chemista, 7th ed., consult Victhods of Analysis of the Association of Official Agricultural Chemista, 7th ed., consult Victhods of Settlery and Green (J. Biol. Chem. 155, 513 (1944)) have suggested a rapid method for the nucrodetermination of arsenic in which the assenic is converted into arsine by electrolysis.

¹¹ Test paper strips are prepared as follows. Saturate filter paper with 5per cent solution of microurie brounds in alcohol allowing the spiper to dry spontaneously while suspended in air Cut into strips 2 5 min. × 12cm and store in a brown glass bottle.

MERCURY

The rapidity of absorption of mercury depends upon a number of conditions such as mode of administration, the nature of the compound and its physical state, the state and condition of the stomach and intestines, the quantity and quality of the food in the stomach, and the state of the circulation of the portal of entrance. There is no definite knowledge as to the form in which it is absorbed. Elimination depends upon the state of the excretory organs, It is eliminated in all the excretions of the body—urine, feces, saliva, sweat, tears, and milk, Elimination begins about two hours after introduction. Depending upon the amount introduced and absorbed and the extent of renal injury, the time required for its complete elimination varies from 24 hours to many weeks.

TESTS FOR MERCURY

- 1. Reinsch's Test. The procedure is carried out in the same manner as for araenic (see above). A piece of arsenic-free copper foil is introduced into the urine acidified with one-fifth its volume of pure hydrochloric acid. The urine is, however, not heated to holling, but warmed to 50° C. or 60° C. and set aside for 12 or preferably 24 hours. Metallic mercury is deposited on the foil as a hright lustrous micro. The foil is then washed with pure water, alcohol, ether, and dried without heat, rolled into a groil, inserted into a glass tube and heated in the same manner as under "Arsenic." The mercury is deposited in the metallic state in the form of globules readily distinguishable with the microscope.
 - 2. Amalgamation Test. A more rapid method than the above is by amalgamation with sine. Add 5 g, of zinc dust to the urine and heat for 15 minutes, sthring continuously. Allow the amalgamated zinc to settle and decant the urine. Then wash by decantation several times with pure water, then with alcohol, and finally with ether and dry in air. Now introduce the dry zinc into a narrow dry glass tube scaled at one end. With the Bunsen huner soften the tube about 2 inches above the zinc and constrict the tube by pulling the ends apart. Introduce a small hit of glass wool or asbestos sufficient to support a small piece of iodine. Introduce the iodine supported by the asbestos at the constriction. Apply heat to the zinc amalgam, and then gently to the region holding the ioding to gently obtaillize it, and immediately reapply heat to the zinc. The mercury volatilizes and, meeting the iodine vapor, unites with it and is deposited as the red iodide of mercury.

LEAD

Lead may be found to the extent of 0.05 mg, or so per liter in the urine of healthy individuals and may be estimated by the method of Fairhall. ¹⁴ It is increased in lead poisoning.

INOSITOL

Inositol, C₄H₄(OH)₄, occasionally occurs in the urine in albuminuria, diabetes mellitus, and diabetes insipidus. It is claimed also that copiou⁵

¹⁰ Fairhall, J. Biol. Chem., 64, 485 (1924), Aub. Fairhall, Minot, and Reznikoff: Medicine, 6, 1 (1925), Millet: J. Biol. Chem., 83, 265 (1925), See also Chapters 23 and 31.

water drinking causes this substance to appear in the urine For further discussion, see Chap 35

TEST FOR INOSITOL

1 Detection of Inositol (Scherer) Acidify the urine with concentrated nitric acid and evaporate nearly to dryness Add a few drops of ammonium hydroxide and a little calcium chloride solution to the moist residue and evaporate the muxture to dryness in the presence of inositol (0 001 g) a bright red color is obtained

For a more satisfactory test, which, however, is more time-consuming, see Salkowski's modification of Scherer's test

FAT

When fat finds its way into the urine through a lesion which hrings some portion of the urinary passages into communication with the lymphatic system, a condition known as *chyluria* is established. The turbid or milky appearance of such urine is due to its content of chyle. This disease is encountered most frequently in tropical countries, but is not entirely unknown in more temperate climates. Albumin is a constant constituent of the urine in chyluria. Upon shaking a chylous urine with ether, the fat is dissolved by the ether and the urine hecomes less turbid or entirely clear Alimentary lipuria may occur following the ingestion of a large amount of fat.

MELANINS

These pigments never occur normally in the urine, but are present under certain pathological conditions, their presence being especially associated with melanotic tumors. Ordinarily the freshly passed urino is clear, but upon exposure to the air the color deepens and may at last he very dark brown or black in color. The pigment is probably present in the form of a chromogen or melanogen, and upon coming into contact with the air oudation occurs, causing the transformation of the melanogen into melania and consequently the darkening of the tirme.

It is claimed that melanuria is proof of the formation of a visceral melanotic growth. In many instances, without doubt, urnes rich in indicin have been wrongly taken as diagnostic proof of melanuria. The pigment melanin is sometimes mistaken for indigo and melanogen for indicin. It is comparatively easy to differentiate between indigo and melanin through the solubility of the former in chloroform.

In rare cases melanin is found in urinary sediment in the form of fine

amorphous granules

TESTS FOR MELANIN

I Ferric Chloride Reaction (son Jaksch Pollak) Add a few drops of ferric chiorida solution to 10 ml of urine in a test tube and note the formation of a gray color Upon the further addition of the chloride a dark precipitate forms, consisting of phosphates and adhering meianin An excess of ferric chloride causes the precipitate to dissolve

This is the most satisfactor; test for the identification of melanin in the urine

¹ Salkowski Z physiol Chem 69 478 (1910)

2 Bromine Test (Zeller) To 50 ml of urine in a small beaker add an equal volume of bromine water in the presence of melanin a yellow precipitate will form and will gradually darken in color, ultimately becoming black

UROROSEIN

Urorosem is a urmary pigment which does not occur preformed in the urine, but is present in the form of a chromogen, which is transformed into the pigment upon treatment with a mineral acid. Herter showed this chromogen to be indolerectic acid.

Indoleacetic acid may be found in urine free or as a compound with glycine Normal urine responds to the uroroscin reaction (see helow) if nitrites are present Pathologically, a positive urorosein reaction is obtained in a variety of diseases, such as pulmonary tuberculosis, typhoid fever, nephritis, stomach disorders, and pellagra The reaction in the urine of pellagra patients has attracted considerable interest because of its possible metabolic and diagnostic significance 36

TEST FOR UROROSEIN

Nitrate Hydrochioric Acid Test (Urorosein Reaction) To 10 ml of urine in a test tube add 2 ml of concentrated hydrochloric acid and a few drops of a 1 per cent solution of potassium nitrite A rose red color indicates uro roseln The chromogen (indolescetic acid) has been changed to uroroseln by oxidation

PORPHYRINS#

Coproporphyrin and uroporphyrin are present in small amount in nor mal urine Both are reddish pigments which may be increased in amount in pathological urine or after administration of quinine tetronal, trional, or sulfonal

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30

Urine: Sediments and Calculi

The sedimentary constituents of the urine may be divided into two classes, viz, organized and unorganized. The sediment is collected by centrifuging the urine at low speed or by allowing it to stand for some time in a conical vessel.

I. UNORGANIZED SEDIMENTS

The more common unorganized sediments consist of ammonium magnesium phosphate ("triple phosphate"), calcium oxalite, calcium phosphate, unc acid, and sodium and ammonium urates Less commonly observed are calcium carbonate, calcium sulfate magnesium phosphate, cystine, leucine, tyrosine, hippuric acid, bibrubin, indigo, xanthine, and melanin

The separation as sediments depends upon the degree of saturation of the urine for these substances, which in turn is influenced by the reaction of the urine Uric acid most commonly separates out from strongly acid urnes, sodium urate from less acid urines Calcium phosphate comes out most commonly in urines more alkaline than pH 6, while calcium oxalate is found in acid, alkaline, and neutral urines. All of these substances are found as seduments in normal urines, and the majority of normal urines show one or more of these separating out on standing Ammonium mag nessum phosphate, ammonium urate, and calcium carhonate precipitate from urmes which have undergone ammoniacal fermentation due to an infection in the urmary tract, and which are honce alkaline in reaction Maslow studied the sediments forming on long standing in carefully preserved specimens of the urines of normal young men Schiments were found in 93 per cent of the urines. Unc acid was found in 17 per cent of cases at an average pH of 55 and sodium urate in a similar number of cases at an average pH of 58 Calcium phosphate was found in 46 per cent of cases with an average pH of 62 and calcium oxalate was found in 71 per cent of cases and at all reactions

Ammonium Magnesium Phosphate (Triple Phosphate") Crystals of triple phosphate are a characteristic constituent of the sediment when alkaline ferments tion of the urine has taken place either before or after being oxided They may even be detected in neutral or slightly acid urine provided the ammonium salts are present in large enough quantity This substance may occur in the sediment in two forms viz prisms and the feather; type The prismatic form of crystals is the one most commonly observed in the sediment the feather; form predominates when the urine is made animonacial with ammonia (see Fig. 226)

The sediment of the urine in such disorders as are accompanied by a retention of urine in the lower urinary tract contains 'triple phosphate' crystals as a characteristic constituent. The crystals are frequently inbundant in the sediment during paraplegia, chronic cystitis, enlarged prostate, and chronic pyelitis.

Calcium Oxalate. Calcium oxalate is found in the urine in the form of at least two distinct types of crystals, viz, the dumbbell and "octahedral" types! (Fig 230) Either form may occur in the sediment of neutral, alkaline, or acid urine, but both forms are found most frequently in urine having an acid reaction. Occasionally, in alkaline urine, the octahedral form is confounded with "triple phosphate" crystals. They may be differentiated from the phosphate crystals by the fact that they are insoluble in acetic acid.

The presence of calcium oxalite in the unne is not of itself a sign of any abnormality, since it is a constituent of normal urine. However variations from the normal occur in certain pathological conditions og parathyroid derangement, urinary calculi, etc.

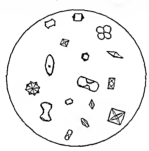


FIG 230 CALCIUM OXALATE (OGDEN)

Calcium Carbonate Calcium carbonate crystals form a typical constituent of the urine of herbivorous animals. They occur less frequently in human urine. The reaction of time containing these crystals is nearly always alkaline, although they may occur in neutral or in slightly acid urine. It generally crystallizes in the form of granules spherules, or dumbbells (Fig. 231). The crystals of calcium carbonate may be differentiated from calcium vashate by the fact that they dissolve in acetic acid with the evolution of carbon dioxide gas.

Calcium Phosphate (Stellar Phosphate). Calcium phosphate may occur in the urine in three forms, viz, amorphous granular, or crystaline. The crystals of calcium phosphate are ordinarily pointed, vedge-shaped formations which may occur as individual crystals or grouped together in more or less regularly formed rosettes (see Fig. 64). Acid sodium turate crystals (Fig. 233) are often mistaken for crystals of calcium phosphate. We may differentiate between these two crystalline forms by the fact that acetic acid will readly dissolve the phosphate, whereas the urate is much less soluble and when it is finally brought into solution and recrystallized, one is frequently enabled to identify une acid crystals which have been formed from the

¹ The so-called octahedral type is strictly speaking a flat, tetragonal dipyramid identical with the mineral weddelite whereas the dumbbell shaped spherulite aggregates correspond to who wellite (Philipsborn Arzikehe Forzek, 9, 391 (1953))

and urate solution. The chineal significance of the occurrence of calcium phosphate crystals in the urinary sediment is similar to that of triple phosphate. (see p. 854)

crystals in the urmary seminates assumed to that are of quite rare occurrence in the Calcium Sulfate Crystals of calcium sulfate are of quite rare occurrence in the sediment of urne. Their presence seems to be limited in general to urnnes which are of a decided acid reaction Ordinarily it crystallizes in the form of long, thin, colorless

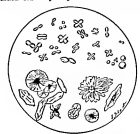


FIG 231 CALCIUM CARBONATE



Fig. 232 Various Forms of Unic Acid 1, Rhombic plates 2 whetstone forms 3, 3 quadrate forms, 4, 5, prolonged into points 6 8 rosettes 7 pointed bundles, 9,

barrel forms precipitated by adding hydrochloric acid to unine needles or prisms (Fig. 225) which may be mistaken for calcium phosphate crystals. There need be no confusion in this respect however, since the suifate crystals are insoluble in acetic acid which reagent readily dissolves the phosphate. Is far as is known, their occurrence as a constituent of urinary sediment is of very little clinical significance.

Uric Acid. Uric acid forms a very common constituent of the sediment of urines which are acid in reaction. It occurs in more varied forms than any of the other crystalline sediments (see Fig. 232), some of the more common varieties of crystals being thombie prisms wedges, dimibbells, whetstones, prismatic rosettes, irregular or hexagonal plates, etc. Crystals of pure uric acid are always coloriess (see Fig. 220), but the form occurring in urinary sediments is impure and under the microscope appears pigmented, the depth of color varying from yellow to a dark reddish brown according to the size and form of the crystal.

The presence of a considerable urise and sediment does not, of necessity, indicate a pathological condution or a urine of increased urise and content, since this substance very often occurs as a sediment in urines whose urise and content is diminished from the normal merely as a result of changes in reaction etc. Pathologically, urise and sediments occur in gout, acute febrile conditions, chronic interstitual nephritis, etc. If the increaseopical examination is not conclusive, urise and may be differentiated



FIG 233 ACID SODIUM URATE

from other crystalline urinary sediments from the fact that it is soluble in alkalics, alkali carbonates, boiling glyccrol, coocentrated sulfure veid, and in certain organic bases such as ethylamine and piperidine. It also responds to the murexide test, Schiff's reaction, and to Folin's phosphotungstic acid reaction (see p. 797).

Urates The wrate sedument may consist of a mixture of the wrates of ammonium, calcium, magnesium, potassium, and sodium. The ammonium wrate may occur in neutral, alkaline, or read urine, whi reas the other forms of urites are confined to the schiments of acid urines. Sodium wrate occurs in sediments more abundantly than the other wrates. There are two sodium wrates, the mono- and the di-, which may be expressed thus.



The so-calkd quadrurate or hemorite have no easterne as chemical units The initiates of calcium, magnesium and potassium are amorphous in character, whereas the urite of aminonium is crystilline bodium urate may be either amorphous or crystilline when crystilline in the constitution of fan-shaped clusters or colorless, prismite needles (Fig. 23) Aminonium urate is ordinarily present in the sediment

in the burthke form of the thorn apple crystal 1e yellow or reddish brown spheres covered with sharp spicules or prisms (1ig 234). The urates are all soluble in hydrochloric acid or acetic acid and their acid solutions yield crystals of uric acid upon standing. They also respond to the murexide test. The clinical significance of urate sediments is very similar to that of uric acid. A considerable sediment of amorphous urates does not necessarily indicate a high uric acid content, but ordinarily signifies a concentrated urine having a very strong sendity.

Cystine Cystine is one of the rarer of the crystalline urinary sediments. It has been claimed that it occurs more often in the urine of men than of women. Cystine crystallizes in the form of thin colorless hexagonal plates (see Fig. 45), which are mosoluble in water, alcohol and acetic acid and soluble in inneral acids alkaless and especially in ammonia. Cystine may be identified by burning it upon platinum foil

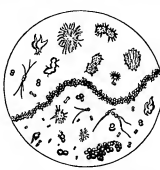


Fig 234 Ammonium Ubates Showing Spherules and Thorn Apple-Shaped Crystals (After Peyer.)

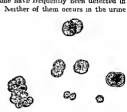
under which condition it does not melt but yields a bluish green flame. For the preparation of cystine see p. 140

Cholesterol Cholesterol crystals have been but rarely detected in urmary sediments. When present they probably areas from a pathological condition of some portion of the urmary tract. Crystals of cholesterol have been found in the sediment in cystitis pyclitis chyluma and nephritis. Ordinarily they occur as large regular and irregular colories transparent plates some of which possess notched corners (see Fig. 101). Frequently instead of occurring in the sediment, cholesterol is found in a film on the surface of the urms.

Hippuric Acid. This is one of the rare sediments of human urine. It deposits under conditions similar to those which govern the formation of uric acid sediments. The crystals which are colories needles or praisa (see Fig. 223) when pure are invariably pigmented in a manner similar to the uric acid crystals when observed in urinary sediment and because of this fact are frequently confounded with the rare forms of uric acid. Hippuric acid may be differentiated from uric acid from the fact that it does not respond to the murexide test and is much more soluble in water and in either. The

detection of crystals of hippuric acid in the urine has very little clinical significance. since its presence in the sediment depends in most instances very greatly upon the nature of the diet. It is particularly prone to occur in the sediment after the ingestion of certain fruits as well as after the ingestion of benzoic acid (see p. 804)

Leucine and Tyrosine Leucine and tyrosine have frequently been detected in the urine, either in solution or as a sediment. Neither of them occurs in the urine ordinardy except in association with the others, i.e., whenever leacine is detected it is more than probable that tyrosine accumpanies it They have been found pathologically in the urine in acute yellow atrophy of the liver in reute phosphorus poisoning, in cirrhosis of the liver, in severe cases of typhoid fever and smallpox, and in leukemia In urmary sedi ments leucine ordinarily crystallizes in characteristic spherical masses which show both radial and concentric striations and arc highly refractive (Fig. 235). Some investigators claim. that these crystals which are ordinarily called leucine are, in reality, generally urates For



235 CRISTALS OF IMPURE LEUCINE (OGDEN)

the crystalline form of pure leucine obtained as a decomposition product of protein see Fig. 42 Tyrosine crystallizes in urmary sediments in the well known sheaf or tuft formation (Fig 44) For other tests on leucme and tyrosine see pp 135 and 138

Billruhin Bilirubin crystallizes in the form of tufts of small needles or in the form of small plates which are ordinarily yellowish red in color (Fig. 100) Pathologically, typical crystals of bilirubin have been found in the urinary sediment in laundice acuto yellow atrophy of the liver, carcinoma of the liver, currhosis of the liver, and in phosphorus poisoning, typhoid fever, and scarlatina

Magnesium Phosphate Magnesium phosphate crystals occur rather infrequently in the sediment of urine which is neutral alkaline or feebly and in reaction. It ordinarrly crystallizes in elongated, highly refractive rhombic plates which are soluble in

acetic acid

Indign Indigo crystals are frequently found in urine which has undergone alkaline fermentation. They result from the breaking down of induxyl sulfates or induxyl glycuronates Ordinarily indigo deposits as dark blue stellate needles or occurs as amorphous particles or broken fragments These crystalline or amorphous forms may occur in the sediment or may form a blue film on the surface of the urine Indigo crystals generally occur in urine which is alkaline in reaction, but they have been detected in acid urine

Xanthine. Yanthine is a constituent of normal urine but is found in the sediment in crystalline form very infrequently, and then only in pathological urine When present in the sediment xanthme generally occurs in the form of whetstone-shaped crystals somewhat similar in form to the whetstone variety of uric acid crystal. They may be differentiated from une acid by the great ease with which they may be brought into solution in dilute ammonia and on applying heat. Nanthine may also form urinary calculi The significance of xanthine in urmary sedurent is not well understood

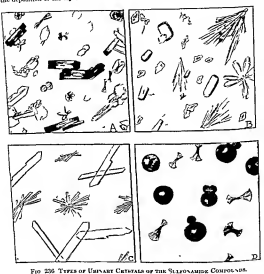
Melanin, Melanin is an extremely rare constituent of urmary sediments. Ordinards in melanuria the niclanin remains in solution, if it separates it is generally held in

suspension as fine amorphous granules

Sulfonamides After administration of the sulfonamides (see p. 657 for discussion of chemical nature) crystalline deposits of the free drug or of its acetylated derivative may be found in the urine Sulfonamide crystals may be recognized by their characteristic appearance (big 230) If the crystals form in the renal tubules during the formation of urine kidnes damage accompanied by hematuria may result. Since the

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compounds producing the crystals are much more soluble in the form of their alkali salts, the maintenance of an alkaline urine during sulfonamide therapy will prevent the deposition of the crystals



A. Sulfathiazole B sulfapyridine C sulfandamide D sulfadiazine Courtesy Kolmer Clinical Diagnosis by Laboratory Examinations D Appleton-Century Co.

II. ORGANIZED SEDIMENTS

Among the more unportant organized sediments are casts of different types, pithelial cells pus cells erythrocytes and microorganisms Cylindroids spermatotoa urethral filaments tusue debris animal parasites fibrin and foreign substances due to contamination are also observed. For detailed discussions of these organized sediments see books on chinical diagnosis or clinical pathology

CALCULI

Urmary calcult, also called concretions or concrements, are solid masses of urmary sediment formed in some part of the urmary tract They vary in shape and size according to their location, the smaller calculi, termed sand or gravel, in general arising from the kidney or the pelvie portion of the kidney, whereas the large calculi are ordinarily formed in the hladder. The condition in which calculi are formed in the human body is termed lithiasis. Very large stones may also form in the kidney. One of these, the so-called staghorn stone, may fill the entire central portion of the kidney. There are two general classes of calculi as regards composition, viz., simple and compound. The simple form is made up of hut a single constituent, whereas the compound type contains two or more individual constituents. The structural plan of most calculi consists of an arrangement of concentric rings about a central nucleus, the number of rings frequently being dependent upon the number of individual constituents which enter into the structure of the calculus. However, layers quite different in macroscopical appearance may be almost identical in composition.

In India and China urinary calculi are most frequently found in children. Up to the middle of the nineteenth century this was true for Europe and America also. But with improvement in diet of more recent years, vesical calculus in children has hecome rare in this country and is now primarily a disease of old age, more than half of persons admitted for operation heing hetween the ages of 50 and 70. Calculi have heen found

in Egyptian tomhs dating as far hack as 4800 B.C.

The etiology of stone in man is still obscure. In rats urinary lithiasis can he regularly produced by diets low in vitamin A. It may he secondary to the epithelial degeneration in the urinary hladder which occurs in vitamin-A deficiency. The calculi contain ammonium magnesium phosphate, calcium carhonate, and calcium hydroxide. The greater frequency of lithiasis in earlier times and the prevalence of stone in children in the Orient may very probably he related to dietary deficiency. Whether vitamin A or other dietary deficiency bears any important relationship to the ctiology of calculus disease in this country at the present time is uncertain.

Urinary calculi have frequently been noted in patients with a variety of bone disorders. In certain of these, associated with hyperparathyroidism, the calculus formation may be related to the markedly increased excretion of calcium in the urine. Immobilization of the patient may also

be a factor.

Stone formation appears often to be secondary to infection in the urinary tract. Clumps of bacteria and epithelial and pus cells may act as foreign-body nuclei for stone formation and the alkaline fermentation frequently associated with infection is favorable to the precipitation of calcium and ammonium magnesium phosphates, which are the most common constituents of secondary calculi. The majority of stones appear, however, to arise in aseptic urines. That stone formation is not commonly due to any metabolic defect is indicated by the fact that most stones are not composed of any one substance but are of a mived type.

Attempts have been made to account for the greater solubility of uric acid and other substances in the urine than in pure water, on the basis of the presence, in the urine, of protective colloids which hinder precipitation. According to this view stone formation may occur when this colloid is altered or diminished in quantity. In support of this view it is stated that the crystal form of calcum ovalate in calculi is not that noted to

form in pure solutions but is similar to that produced when precipitation occurs in colloidal solutions. It is possible that precipitation of colloidal material with the crystalloids does conduce to the formation of a concrement rather than a more precipitate of the latter

According to the investitation of Meyer however, the growth of a concrement depends solely on the degree of saturation of the urne for the crystalline constituents and such precipitation of the crystalline constituents occurs in just the same way in urne as in pure aqueous salt solutions. In determining the nature of the sediment or calculus formed the pH is a controlling factor. Urne acid tends to precipitate out of normal urnes of average composition when the acidity becomes high (in the neighborhood of pH o). Ut pH 6 mixed calculi of urice and sodium urate and calculin oxalate and phosphate will tend to form A pH 7 calcium phosphate calculi would tend to form and between pH 7 and 8 with the urne ammoniaced mixed calculin of calcium phosphate magnesium ammonium phosphate and ammonium urate may form which may he firm if precipitation occurs slowly. It pH values above 8 the rapid precipitation would tend to produce soft stones containing calcium carbonate ammonium magnesium phosphate and ammonium urate.

Aside from the question of protective colloids certain of the stoneforming constituents may under certain conditions themselves exist in part in colloidal solution. Thus Hammarsten found that lithium and potassium urates gave true solutions at 37°. Sodium and ammonium irites in saturated solution are present partly in colloidal form. It the average chloride concentration of the urine the solubility products of the latter may be increased 300 per cent. Urates and urie acid in the presence of chlorides thus have a strong tendency to form supersaturated solutions.

According to Stillman? out of old urnary calcul submitted for routine examination over a period of five to six years at the New York Hospital about 11 per cent consisted largely of calcium carbonate calcium phosphate or triple phosphate eitherations or in mixture about 49 per cent were largely calcium oxidate with some instances of admixture with phosphate and about 6 per cent were largely urice acid Cysture circul occurred to the extent of 0.8 per cent. These examinations refer to the principal constituent of the calculus no attempt was made to isolate a nucleus and analyze it. Ultamann has reported that in 3-b cases of urnary constituent of the calculus and analyze it.

(hap 3	50		URINE SEDIMENTS AND CALCULI						80
	Uric acid	Ammonum urate	\anthue	Cystine	Urostealth	Fibru	Caleum earbonate	Calcum oxalate	Bonc earth (magne- sum and calcaum phosphate)	Triple phosphate, (mixed with unknown amount of earthy phosphate)
	No notecable ammonta reaction	Strong ammonia reaction	e test Tho p	Hame pale blue burns a short time Pecular sharp odor. The powder dis solves in annona and six-sided plates separate on the spontaneous evaporation of the annona	Hame yellow pale contanuous Odor of rean or shellae on burning Powder lubie in alcohol and ether	Thane yellow containous Odor of burnt feathers Insoluble in alcolol and ether Soluble in potassium hydroxide with heat Precipitated therefrom by accid with generation of hydrogen sullido	I ffervesces	Liferytaces	The powder and or HGT This solution gives an phosphatt, mostering amorphous preceptate with animona	Abundant anunonn The powder Triple phosphate's desolves an acete acad or HGI This (mack with unknown soutton grees a crystalline preceptrate announ of earthy with anmouns
	The powder when treated with NOH gives		Does not give murexi effervescence. The dried red on warming.	Hame pale blue burn solves in anmonia and ration of the ammonia	I lame yellow pale continged	Hame yellow contanuous Odor of burnt feat ether Soluble in potassium hydroxide with he acetic acid with generation of hydrogen sullido		The powler gently heated then treated with HG!		
	The powder	The powder gives the marcyide test		I lame pa solves in a ration of th					Does not effervesce	
	Without			With flame			The powder when treated with HCJ			
				Бигпя			Does not			
		On heating the powder on platmun foll, it								

Butt, Hauser, and Seifter² have proposed a therapy for renal lithiasis which embraces a pronounced increase in the urinary protective colloids through the subcutaneous injection of hydluronidase in isotonic sodium chloride solution (see "Connective Tissue," p 243)

VARIETIES OF CALCULI

Urie Acid and Urate Calcuil Urie acid and urates constitute the nuclei of a large proportion of urmary concretions but stones which consist chiefly of uric acid or urates are found in less than 1 out of 10 cases Such stones are always colored the tint varying from a pale yellow to a brownish red. The surface of such calculi is generally smooth but it may be rough and uneven

Phosphatic Calcull Ordinarily these concretions consist principally of triple phosphate and other phosphates of the alkaline earths with very frequent admixtures of urates and oxalates. The surface of such calculi is generally rough but may occa sionally be rather smooth. The calculi are somewhat variable in color, exhibiting gray white or yellow tints under different conditions. When composed of earthy phosphates the calculi are characterized by their friability

Calcium Ozalate Calculi These calcult are quite hard and are rather difficult to crush They ordinarily occur in two general forms viz the small smooth concretion which is characterized as the hemp-seed calculus and the medium-sized or large stone possessing an extremely uneven surface which is generally classed as a mulberry calculus. This roughened surface of the latter form of calculus is due in many in stances to protruding calcium oxalate erystols of the octahedral type

Calcium Carbonate Calculi Calcium carbonate concretions are quite common in herbivorous animals, but are of exceedingly rare occurrence in man. They are gen erally small white or grayish calculi spherical in form and possess a hard smooth surface These considerations apply to stones consisting largely if not entirely of cal cium carbonate. Mixed calculi which contain smaller but readily demonstrable amounts of calcium carbonate are rather common in man

Cystine Calculi Cystine calculi are of very rare occurrence the incidence being us sally less than 1 per cent. Ordinarily they occur as small smooth, oval or cylindrical concretions which are white or yellow in color and of a rather soft consistency

Xanthine Calculi This form of calculus is somewhat more rare than the cystine type The color may vary from white to brownish yellow Very often uric acid and urates are associated with xanthine in this type of calculus. Upon rubbing a xanthine calculus it has the property of assuming a waxlike appearance

Urostealith Calculi This form of calculus is extremely rare Such concretions are composed principally of fat and fatty acid. When moust they are soft and elastic but when dried they become brittle. Urostealiths are generally light in color

Fibrin Calcull Fibrin calculi are produced in the process of blood coagulation within the urmary tract. They frequently occur as nuclei of other forms of calculus They are rarely found

Cholesterol Calculi This is an extremely rare form of calculus somewhat resem I hng the cystme type

indigo Calculi In ligo calculi are extremely rare

The scheine proposed by Heller and given on p 863 will be found of much assistance in the chemical examination of urinary calcul-

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31

Urine: Quantitative Analysis

In analyzing a normal or pathological urine quantitatively for any of its constituents, it is particularly necessary that the complete and exact 24-hour sample be obtained For directions with regard to the collection and preservation of urine for analysis, see Chapter 27 "General Characteristics of Normal and Pathological Urine" Methods for the determination of the specific gravity of the urine are also there described Before any urine is taken for analysis its total volume should be measured, using a large graduated cylinder, and this volume is thereafter taken as a basis for the calculations of the daily output of the individual constituents determined

PREPARATION OF STANDARD ACID AND ALKALI SOLUTIONS

Principle. Many of the quantitative methods used in physiological ebemistry are volumetric or titration procedures For these methods solutions of accurately known strength called standard solutions are needed Their strength is usually expressed in terms of normality A normal solution is one which in 1000 ml contains 1 g of replaceable hydrogen or its equivalent Thus, to make 1000 ml of a normal solution of hydrochloric acid (HCl), we would need 36 5 g of this acid containing 1 g of replaceable hydrogen This we derive from the fact that the atomic weight of Cl is 35 5 and of H is 1, so that the molecular weight of HCl is 36 5, and cach 36 5 g of this acid contains 1 g of replaceable by drogen Sulfuric acid (H-SO₄) has a molecular weight of 2 + 32 + 64 = 98, but 98 g of sulfuric acid contains 2 g of replaceable hydrogen Therefore, to prepare a normal solution of this acid, we must use one-half of 98 or 49 g of sulfuric acid (containing 1 g of hydrogen) for 1000 ml of normal solution. Oxalic acid (H₂C₂O₄ + 2H₂O) has a molecular weight of 2+24+64+36=126 It also is a dihasic acid so we must use only one-half of 126 or 63 g of oxahe acid in making a liter of normal solution

A normal alkali solution is exactly equivalent to a normal acid solution, i.e., I liter of the alkali will neutralize I liter of the acid. According to the reaction of neutralization, therefore, the 36 5 g of HCl in a liter of this normal acid will require 40 0 g of sodium hydroxide to neutralize it, and I liter of normal sodium hydroxide must contain 10 0 g of the alkali

$$11C1 + \text{VaOH} \rightarrow \text{VaCl} + 11_2O$$

(36.5) (40.0) (58.5) (18)

[&]quot; we table of atomic weights. Appendix

Having prepared solutions of acid and alkali of definitely known strength, it is then possible to determine the strength of any unknown acid or alkali hy finding out much much of these standard solutions is required to neutralize a definite volume of the unknown solution

In order to tell when the unknown solution has been exactly neutralized, we titrate in the presence of a small amount of one of a class of substances called indicators. An indicator is a substance which undergoes a sharp color change at a particular range of hydrogen-ion concentration, and this color change indicates the end-point of the titration

When strong acids (as HCl) are being titrated with strong alkalies (as NaOH), almost any one of the common indicators is satisfactory. If weak acids (acetic acid) or weak bases (as ammonia) are being titrated, it is necessary to be very careful in the choice of an indicator as all indicators are not equally satisfactory in these instances * (See p 39)

Preparation of 0.1 N Oxolic Acid Solution. Weigh accurately a watch glass or a piece of glazed paper. Then add to the weights on the balance pan 3.1512 g. With a spatula transfer to the watch glass enough pure oxalic acid in the form of clear crystals to counterhalance exactly the weights in the opposite pan Transfer completely to a 250-ml, heaker, using a little water for rinsing purposes Add about 150 ml, of distilled water and stir with a glass rod until dissolved, warming gently if necessary Transfer every particle of this solution to a clean 500-ml volumetric flask, rinsing rod and heaker several times with distilled water Hold under the tap until cooled to room temperature. Then add distilled water until the bottom of the meniscus is level with the mark on the neck of the flask (the lower mark if there are two) Insert a stopper and mix thoroughly by inverting the flask again and again. Transfer to a clean dry bottle Label. This solution will not keep indefinitely and is to be used only in the standardization of 0 1 N alkali.

Preparotion of 0 I N Sodium Hydroxide Solution.

(a) PREPARATION OF CONCENTRATED CARBONATE FREE SOMEM HYDROUGE SOLUTION Shake up about 110 g, of best quality NaOH with 100 ml of distilled water in a 300-ml. Erlenmeyer flask (pyrex) to make a saturated solution. Stopper and allow to stand for a couple of days or until the sodium carbonate settles to the bottom, leaving a clear solution of NaOH practically free from Carbonate.

(b) PREPARATION OF A STANDARD SOURM HYDROXIDE SOLUTION Measure out 6 3 ml. of the saturated NaOII solution from a buret into a 1-liter flask, Add 750 ml. of distilled water and mix thoroughly. Clean a buret by allowing it to stand filled with cleaning mixture (sodium dichromate and sulfuric acid) for a few minutes or longer if necessary. Empty, rines several times with tap water, finally with distilled water, and allow to drain. Introduce a few ml. of the NaOII solution, and invert a couple of times to rinse the buret, discriding this NaOII. Repeat this process at least twice more. Then fill the buret with the alkall solution, making sure that the tip contains no air bubbles, and run out solution until the bottom of the meniscus is exactly at 0.

Into a clean Erienmeyer flask. (150 to 250 ml.) now Introduce 25 ml. of 0 1 \(\) oxalle acid solution measured from an accurate, clean pipet, previously rinsed by means of a little of the acid solution drawn up into it. Allow the pipet to drain about 15 seconds against the side of the flask. Add 2 to 3 drops of a 1 per cent alcoholic solution of phenolphthalein.

For further consideration of indicators see pp 38 and 378

ow run in NaOli solution from the buret, rotating the flask. Ten mi. .. n be added quite rapidly, then add more slowly, and finally drop by drop until the last drop changes the color of the solution permanently throughout to a definite pink. Take the buret reading, estimating as closely as possible to the second decimal place. Repeat the titration until two closely agreeing dupiicate readings are obtained, then average the two readings for calculation.

Calculate the strength of the NaOII solution. Divide 25 (the number of mi. of 0.1 N oxalic acid used) by the buret reading to obtain the strength of the NaOll in terms of 0.1 N solution. Then multiply by 0.1 to obtain the normality. For example, if 15.67 ml. were required: 25 - 15.67 = 1.595;

1.595 × 0.1 = 0.1595 N.

(c) PREPARATION OF THE 91 N NAOH SOLATION Calculate how much of the standard NaOll solution just prepared will be required to make I liter of 0.1 N solution. To do this divide 1000 ml, by the strength of the NaOII in terms of 0.1 N solution. Thus to the example cited above: 1000 - 1.595 = 626.9 ml. required. Measure out the exact amount of alkali required (using the buret, pipet, and volumetric flasks) into a 1000-ml. flask. Dilute with distilled water exactly to the mark. Mix very thoroughly and transfer to a clean, dry bottle with a rubber (not glass) stopper. Check the strength of the solution by again titrating 25-ml. portions of oxalic acid solution.

Preparation of 0.1 N Hydrochloric Acid. Concentrated hydrochloric acid is about 12 N or 44 per cent HCl weight in volume. Approximately 0.1 N HCl may, therefore, be prepared by diluting 9 ml. of the concentrated acid to I liter In a volumetric flask. This must be standardized by titration with 0.1 N alkali, using aiizario red or methyl red as an indicator.

Or introduce into a 1-liter flask 12 ml. of coocentrated IICl and 750 ml. of distilled water. Mix well and titrate 10, 15, or 25 ml. portions of the acid solution with 0.1 N NaOII, using alizarin as an indicator. Dividing the number of ml. of 0.1 N NaOH required by the number of ml of acid used gives the strength of the 11Cl in terms of 0.1 N solution. Dividing 1000 by this quotient gives the number of mi, of 11Cl solution to be measured into a volumetric flask and made up to 1000 ml.

This diluted solution will be 0.1 N HCl It should be mixed thoroughly and 25-ml, portions of it checked by tstration with the 0.1 N NaOll.4

Standard acid and alkali solutions are best kept in paraffin-lined bottles The acid solution is the more permanent of the two; and standard sulfurio acid solutions are more permanent than hydrochloric acid solutions Alkalı solutions must be protected from the carbonic acid of the air, the solution being best drawn over into the buret by means of a siphon tube leading from the top of the buret to the interior of the alkali bottle The

If a very high degree of accuracy is desired, the alkali may be checked against pure acid potassium phthalate (mol wt 204 139) Dodge J Ind Eng Chem 7, 29 (1915), J Am-Chem. Soc , 42, 1655 (1920)

The acid solution may be standardized directly in the following manner Introduce a platinum dish containing very pure sodium bicarbonate or the highest grade anhydrous sodium carbonate into a hot-air over previously heated to 200° C. Raise the temperature to 270° to 280°, but not above 300° C. Heat for half an hour allow to cool in a desicator but while still a little warm, transfer to a glass-stoppered weighing bottle Weigh out rapidly 0 1- to 0 2-g. portions of the sodium earbonate, dissolve in about 50 ml of water in an Erlenmeyer flask, and titrate, using methyl orange as an indicator One hundred ml. of 0 l N acid is equivalent to 0.530 g. of dried sodium carbonate

air inlet through the stopper of the bottle should be guarded by a tuke containing soda lime

Preparation of Stondord Hydrochloric Acid from a Constant Boiling Solution (Method of Hulett and Bonner) Procedure Make up by hydrometer about 200 ml of HCl of specific gravity about 110 Distil off about three-fourths of the hauld and discard Then collect about 25 ml of the constant-boiling distillate Weigh 18 017 g of distillate using a capillary piper for final adjustment (or measure out 16 442 ml at 25° C) Dilute to 1 liter This is a 0 1 N solution of HCl. The figures given are for a barometric pressure of 760 mm At 770 mm use 18 039 g, at 750 mm use 17 996 g, at 740 mm use 17 975 g and at 730 mm use 17 953 g. The composition of the distillate should not vary more than 1 part in 10.000 from the figures given

TITRATABLE ACIDITY OF LIRINE

Introduction. The titratable acidity of the urine, as determined by the Γ ohn method described here, is expressed in terms of the amount of standard alkali necessary to bring the urine from its original pH to the phenolphthalein end point, around pH 85 or 9 In normal urine, the "acid" iterated consists almost entirely of the acid phosphate ion, $H_2PO_4^-$, the reaction being as follows

$$H_2PO_7 + OH = HPO_7 + H_2O$$

Small amounts of titratable acidity may be contributed normally by acid organic salts (acid urate, acid oxalate, etc.) In certain conditions such as severe ketosis sigmificant amounts of free \(\textit{p}\) hydroxybutyric acid, for example, may be present in the urine and contribute to the titratable acidity. In general, however, monobasic organic acids are found in the urine largely if not entirely as their salts, and in this form contribute hitle or nothing to the total titratable acidity. To evaluate the significance of the exerction of organic acids on acid base balance, they may be determined as such (see "Determination of Organic Acids," p. 371) or the urinary ammonia content determined as an index of base replacement (see p. 883).

The Poin method is theoretically unsound as an exact measure of acid excretion relative to the normal acid base balance in the body, and cannot be used for such purposes. A more accurate picture is obtained if the urine is titrated back to the pH of the blood (pH 74) rather than to pH 9, since it is the difference between the pH of blood and urine which reflects the excretion of acid or base in excess of the amounts normally required for acid base balance. A method for determining titratable acidity by titrating urine to pH 74 against a color standard is described by Henderson and Palmer, a pH meter may also be used in the Polin method, urines even more alkaline than the blood may show a positive "fitratable acidity," the quantitative significance of which is obscure. Despite these limitations, the method is simple and has been widely used, especially for comparative purposes and in conjunction with

³ Henderson and Palmer J Biol Chem. 17 305 (1914)

the determination of urmary ammonia as an aid in estimating the seventy of acidosis, as described by Fitz and Van Slyke **

Folin's Method, Principle. The urine is titrated with standard sodium hydroxide solution, using phenolphthali in as an indicator Potassium oxidate is added to pre-cipitate the calcium which would otherwise interfire with the end point due to the precipitation of calcium phosphate on neutralization of the urine

Procedure. Place 25 ml. of urine in a 200-ml. Erlenmeyer flask and add 5 grams of finely pulverlzed potassium oxalate and 1 to 2 drops of a 1 per cent phenoiphthalein solution to the fluid. Shake the mixture vigorously for to 2 minutes and titrate it immediately with 0.1 N sodium hydroxide until a faint hut unmistakable pink remains permanent on further shaling For more accurate results, particularly with deeply colored urine, compare during the titration against a control sample of urine plus oxalate, until there is a distinct difference in color in the titrated sample. The control sample may then he used for a duplicate estimation, titrating it to color match against the first sample. Take the buret reading and calculate the titratable acidity of the urine under examination.

CALCULATION If y represents the number of milliliters of 0 1 N sodium hydroxide used and y' represents the volume of urine exercted in 24 hours, the total acidity of the 24-hour urine specimen (x) may be calculated by means of the following proportion

25/y = y'/x (titratable acidity of 24-hour urine expressed in ml of 0.1 NaOII)

Interpretation. The titratable acidity of the urine, expressed in ml of 0 1 N alkali required to neutralize the 22-hour output by the method described, varies ordinarily from 200 to 500 under normal conditions with an average of perhaps 350 It is dependent almost entirely upon the diet, heing low on a vegetable (hase-forming) diet and high on a diet containing much meat, milk, cheese, rice, whole wheat products, etc (acid-forming foods) On the administration of 15 g of sodium bicarhonate it may foom to 100, the ingestion of much acid-forming food may increase it to 600 In fasting it may rise in a few days to 800 Acidities of less than 250 usually indicate a true alkalinity of the urine inasmuch as phenolphthalen changes at a pH significantly more alkaline than that of the blood, as discussed above Samples of urine collected shortly after a meal may be alkaline due to the so-called "alkaline the".

Bacterial decomposition of the urea of the urine occurring in the urinary tract will increase the amount of ammonia and decrease the aeduty of the urine. The same change usually occurs in urine left in coatact with the air. The aedity of the urine is increased in acidosis and cardiorenal and certain other disorders. The aedity of the urine may be somewhat increased by administration of mineral acids, acid phosphates,

Fits and Van Slyke J Biol Chem 30, 339 (1917) Van Slyke J Biol Chem 33, 271 (1918)
 Barnett J Biol Chem 33 267 (1918)

⁴ Under the heading interpretation there will be found, in connection with the various quantitative methods which follow brief notes as to the possible nignificance of the results obtained For further discussions see Chapters 27 33 and 34 General references will be found listed at the end of the chapter particularly with reference to climical aspects.

or ammonium chloride, hut it is much more difficult to increase than to decrease this acidity

Determination of Organic Acids (Method of Von Slyke and Polmer) ⁷
Principle Carbonates and phosphates are preemptated and the filtrate titrated with
acid from pH S to pH 27 In the procedure described, the indicators phenolphthalen
and tropcolm OO are used to define the pH range a pH meter may also be used
Variations in the technique are required in the albuminous or high bicarbonateurines

Procedure Mix 100 ml of urine with 2g of finely powdered calclum hydroxide and stir occasionally for 15 minutes Filter Carbonates and phosphates are removed Transfer 25 ml of filtrate to a 125 to 150 ml test tube, add 0 5 ml of 1 per cent phenolphtbalein and 0 2 N HCl from a buret until pluk color just disappears This amount need not be measured The plt is now about 8 Add 5 ml of 0 02 per cent tropeolin OO* little by little with sturing to prevent precipitation of the dye Titrate with 0 2 N HCl until the red color matches that of a standard (plt 27) made from 0 6 ml 0 2 N HCl, 5 ml of tropeolin OO solution and water to make 60 ml When the end point is nearly reached dilute unknown to 60 ml before completing the titration

To avoid the buffering effect of protein in albuminous urine, acidify with a few drops of concentrated HCI, boil, and filter before proceeding with the addition of calcium bydroxide as described above Similar acidification and aeration by pouring will sufficiently reduce the CO; content of high bicar bonate urines (pil) over 7)

CALCULATION Subtract from ml 0.2 N HCl the amount (usually 0.7 ml) of the acid required to titrate a control tube of water between the same limits. Multiply the difference by 2 to get results in terms of 0.1 N acid (the usual basis) and by 1000/2a to get the value for 1000 ml of urine

Interpretation. The titration includes organic acids, creatine, creating, and a small amount of amino acids. Normally the excretion for 24 hours corresponds to 400 to 750 ml of 0.1 N HCl or about 8 ml per kg of body weight. In diabetic acidosis values from 20 to 180 ml per kg have been observed. In diabetic this titration approximates closely the amount of β -hydroxybutyric and acetoacetic acids in the urine and may be substituted for the determination of these substances for clinical purposes

HYDROGEN-ION CONCENTRATION OR TRUE ACIDITY

Colorimetric Method Principle The reaction of the unine is determined by matching the colors produced when pH indicators are added respectively to the diluted unine and to the standard solutions of known reaction. This method is subject to salt and dilution errors but is sufficiently accurate for most clinical purposes.

Procedure Preparation of Standard Buffer Solutions. The Sørensen phosephiate buffer standards, together with the Walpole acctate standards, consatisfactorily the range of pH of urine. The acctate buffers are prepared by mlxing 0 1 N acetic acid and 0 1 N sodium acctate in the following proportions

¹ Yan Sble and Palmer J Biol Chem. 4: 56" (19°0) Palmer J Biol. Chem. 48, 245 (1996) Reventing to Widmark and Liquidpere (Blochem Z 216 1 (19°9)) the Cad used in this method carries down from 7 to 10 per cent of the ether-soluble acids of urine and thus Bive salgith by results.

^{*} In certain urines this indicator fades under acid con litions. Bron phenol like may be substituted or preferably a pH meter may be used

pH	0,1 N Acetic Acid	0.1 N Sodium Acelal	
1	ml.	ml.	
.6	185	15	
.8	176	24	
	164	36	
.0	147	53	
.2	126	74	
	102	98	
8	80	120	
	59	141	
5.0 5.2	42	158	
	29	171	
5.6	19	181	

The phosphate buffers are prepared by mixing M/15 disodium phosphate and M/15 potassium acid phosphate in the proportions given in the table below.

p∐	M/15 Na ₁ HPO ₄	M/15 KH PO	
	ml.	ml.	
5.4	3.0	97 0	
5.6	50	95 0	
5.8	7.8	92.2	
6.0	12 0	88 0	
62	18 5	81.5	
6 4	26 5	73 5	
00	37.5	62.5	
68	50 0	50 O	
70	61 1	38 9	
7 2	71 5	28 5	
7 4	80 4	19 6	
76	86 8	13.2	
7 8	91 4	8.6	
2 11	1 915	7 22	

ards, introduce 8 ml of recently boiled distilled water and 10 drops (or 0.5 ml) of a particular indicator solution, say bromocresol purple. To this diluted indicator solution now add a layer of mineral oil and then introduce, without exposure to alr, 2 ml of urine. Stir gently and match with indicator standards in a comparator block. A control tube containing the urine diluted 1.5 should be used in making the comparison, as described in the hydrogenion determination of blood (see p. 699).

If the urine is too acid or alkalline to come within the range of the indicator selected, repeat, using a more suitable indicator, until a satisfactory one is found

CALCULATION The colorimetric reading made on the diluted urms at room tempers ure may be approximately corrected to give the actual pH at 38° C by subtracting 0.2 pH. The latter factor is however not constant. Hastings Sendroy and Robson warm the diluted urms to 38° C before reading and correct only for the dilution by subtracting 0.1 pH. This gives results which are claimed to be within 0.05 pH of the electrometric pH of the undiluted urns at 38° C.

Interpretation. Tho pH of urme may vary over the extreme range of pH 48 to 80, normal pH values he hetween about 55 and 80, with a mean value of about 60 In most pathological conditions the mean pH is lowered (increased acidity) The factors influencing urmary pH are similar to those influencing the titratable acidity of the urine (diets containing acid- and base-forming foods and the administration of acid or alkali), the two are not necessarily related quantitatively honever. Thus a dilute urme and a concentrated urme may have the same pH, but quite different titratable aciditics. The pH may be considered as an intensity factor, while the titratable acidity is a capacity factor. The determination of urmary pH is of importance clinically largely in relation to the precipitation of insoluble material from the urine and the possible formation of urmary calcult For example, in acid urine, urates and certain sulfonamides may precipitate out, prophylaxis entails the maintenance of an alkaline urine in which these substances are more soluble. The reverse is true for phosphate calculi, which form in alkaline urine but are soluble in acid urine Unnary pH may be lowered by the administration of am monum chloride or raised by sodium bicarbonate

Electrometric Method. The colorimetric determination of urmary pH has been largely replaced for research purposes by the use of the glass-electrode pH meter (see p 48) More accurate results are obtained, and

color or turbidity of the urine does not influence results

TOTAL SOLIDS

Drying Method Place 5 ml of urine in a weighed shallow dish, acidify very slightly with acetic acid (1 to 3 drops), and dry it in vacuo in the presence of sulfuric acid to constant weight Calculate the percentage of solids in the urine sample and the total solids for the 24 hour period

Interpretation The average excretion of total solids by a normal adult man is about 70 g. It is largely dependent upon the protein and salts of the diet. It may be decreased in severe nephritis due to impaired exerction, and greatly increased in diabetes with high sugar climination.

Practically all the methods whose technique includes evaporation at an

increased temperature, either under atmospheric conditions or in racio, are attended with error.

Shackell's method which entails the vacuum descention of the frozen sample is extremely satisfactory and should be used where the greatest accuracy is desired.

Long's coefficient (2.6) was at one time used to compute the solids content of time. The second and third decimal figures of the specific gravity were multiplied by 2.6 to give the number of grams of solid matter in 1 liter of time. This procedure is now ob-ofets.

rotatory shaking and distif the mixture until its volume has diminished about one-haif. Titrate the partly neutralized 6.1 N sulfuric acid solution by means of 0.1 N sodium hydroxide, and calculate the content of nitrogen of the urine examined.

CALCULATION Subtract the number of ml of 0 1 N sodium hydroxide used m the titration from the number of ml of 0 1 N sulfuric acid taken. The remainder is equivalent to the number of ml of 0 1 N sulfuric acid neutralized by the aminonia of the urine. One ml of 0 1 N sulfuric acid is equivalent to 0.0014 g of nitrogen. Therefore, if y represents the volume of urine used in the determination, and y' the number of ml of 0 1 N sulfuric acid neutralized by the aminonia of the urine, we have the following proportion.

 $y/100 = (y' \times 0.0014)/x$ in which x equals g. nitrogen per 100 ml. urine examined

Calculate the quantity of nitrogen in the 24-hour urine specimen

Interpretation. An adult of medium size on a mixed diet will usually excrete 12 to 18 g of mitrogen per day in the urine. On a low protein diet the urinary nitrogen may drop to as low as 5 g or so per day, on a high-protein diet, it may rise to 22 to 25 g or even more. The total mitrogen determination includes the nitrogen from all the introgenous constituents of the urino, under ordinary circumstances the compound urea contributes about 80 per cent or more of the total mitrogen, the remainder being distributed between the various other nitrogenous constituents present (see below). In a normal adult, the total nitrogen of the urine and of the feces will often be almost exactly equal to the total nitrogen of tho diet. Such a condition is called "nitrogen cquilibrium". The feces usually contain very little nitrogen, it is a customary approximation in metabolic studies to assume that the fecal nitrogen will be 10 per cent of the urinary mitrogen.

Calculation of Percentage Nitrogen Distribution In modern metabolism studies where the various forms of introgen are determined, in addition to the total introgen as yielded by the Kjeldahl method, it is customary to indicate what portion of the total introgen was present in the form of each of the individual introgenous constituents. These perceutage values are secured by dividing the weight (grains) of introgen excreted for the day in the form of each individual introgenous constituent by the weight of the total introgen output for the same period. For example, if the total introgen excretions of introgen in the forms of ammonia and creatinnes are 0.271 g and 0.639 g respectively, the percentage distribution for these forms of introgen would be calculated as follows:

```
8 520 g urea mirrogen — 9 814 g total mirrogen = 81 3 per cent
0 271 g ammonia mirrogen — 9 814 g total mirrogen = 2 7 per cent
0 639 g creatinine-mirrogen — 9 814 g total mirrogen = 6 5 per cent
```

NITROGEN PARTITION IN URINES CONTAINING ALBUMIN If the uring to be tested contains albumin, this must be removed before an attempt at a introgen partition is made. This may be done by heating to boiling, accidifying with acetic acid to coagulate the protein, filtering, and making up the filtrate to the original volume of the urine. If very small amounts

of albumin are present, removal is more difficult. In these c uses the use of aluminum hydroxide cream is recommended since it removes none of the nitrogenous constituents of normal urine

Procedure One liter of urine (containing not over 1 per cent of albumin) is mixed with 1 liter of aluminum hydroxide cream¹⁴ and filtered

2 Folin-Farmer Micro-Kjeldahl Method ¹³ Principle. This method belongs with the so-called imerochemical methods massively as it is adapted to the determination of amounts of nitrogen in the neighborhood of 1 mg while in the ordinary Kjeldahl procedure 30 to 100 mg of nitrogen is generally manipulated. One ml of diluted urine is decomposed with sulfure acid, as in the Kjeldahl method, the ammonia formed is set free by the addition of alkah, and carried over into an acid solution by means of a current of air. The ammonia solution is then treated with \(\cdot \setminus \) resgers and the color produced compared with that of a standard solution of an ammonium sall treated in the same way. The reaction is

$$NII_4OII + 2(KI)_2IIgI_2 + 3KOII \rightarrow NIIg_4I + 7KI + 1II_4O$$

The solution of dimercuric ammonium iodide is colloided. The conditions of formation must be carefully controlled to obtain uniformity and prevent pracipitation. Chlorides decrease the color formation.

Procedure Introduce 5 ml of urine into a 50-ml volumetric flask if the specific gravity of the urine is over 1 018, or into a 25-ml flask if the specific gravity is less than 1 018 " Fill the flask to the mark with distilled water and invert it several times in order to guarantee thorough mixing Transfer 1 ml of the diluted urine to a large (20 to 25 mm by 200 mm) pyrex glass test tube To this add 1 ml of concentrated sulfuric acid, 1 g of potassium sulfate, I drop of 5 per cent copper sulfate solution, and a small, clean, quartz pebble or glass bead (The pebble or bead is added to prevent bumping) Boil the mixture over a microburner for about 6 minutes. I e , about 2 minutes after the mixture has become clear and light green or almost colorless Allow to cool until the digestion mixture begins to become viscous This ordinarily takes about 3 minutes, but in any event the mixture must not be permitted to solidly Add about 6 ml of water (a few drops at a time, at first, then more rapidly) to prevent solidification To this acid solution add an excess of sodium hydroxide solution (3 ml of a saturated solution are sufficient), adding the alkali down the side of the tube so that it does not mix with the acid solution but forms a layer beneath tt Connect the tube with a suitable receiver in an aeration train (see Ftgs 238 and 239) placing in the receiver " about 20 ml of dilute (approximately 0 02 N) sulfuric acid Start the air current (which roixes the alkali and acid in the digestion tube) and aspirate the

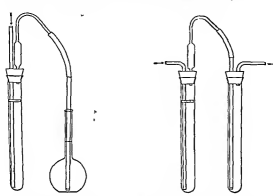
¹⁴ See Appendix

¹⁹ Folin and Farmer J Biol Chem 11 493 (1912)

¹⁹ The purpose is to dilute the urine so that 1 ml of the diluted fluid shall contain 0.75 to 1.5 mg of nitrogen

If For a angle analyses a volumetric flack is used as receiver and compressed as its used for aspuration. For serial analyses a receiving table similar to the digestion tube and fitted with a two-lude rubber stopper carrier as long saled and a short outle used there compressed are or suction obtained from a water pump and applied at the end of the train may be used. The entering are from a water pump and applied at the end of the train may be used. The entering are from two washed by passage times that washed bottle containing duluted 1.10 yellufure and the robe by creaming duluted 1.10 yellufure and the results of the present As many pairs of tubes as desired may be connected a summon which the be present cubber tubing used for con nections see p 5.55 if station times. For middle of cleaning from the complete decreases the rate of air passage to a slow stream and start at the end farthest from the pump to avoid back pressure.

liberated ammona into the receiving solution. The air current should be started slowly and should be only moderately rapid for the first 2 minutes, but at the end of this two-minute period the current should be run at its maximum speed for an interval of 8 minutes, or until all the ammonia has been driven over. The time required for complete aspiration of the ammonia will differ under different conditions and should be established by trial. ¹⁰



Figs 238 (left) and 239 (right) Forms of Aeration Apparatus (Folin and Farmer)

When all the ammonia has been aerated over, disconnect the receiver, dilute the contents to about 60 ml with ammonia-free water in a 100 ml volumetric flask (if such a flask has not itself been used as receiver), and dilute similarly 1 mg of nitrogen in the form of ammonium sulfate" in a second volumetric flask. Nessierize both solutions as nearly as possible at the same time with 10 ml of Nessler solution measured in a graduated cylinder and diluted, immediately before using, with about 20 ml of am monla-free water to avoid turbldity immediately fill the two flasks to the mark with ammonia-free water, mix well and allow to stand 5 minutes before

¹⁸ Run a trial analysis and have several receivers ready. After S minutes aspiration remove the first receiver and replace by a fresh one. Repeat at everal later time intervals. Treat the contents of each receiver with Needler solution as described above for an analysis and estimate from the color reaction the aeration time which gives complete transference of animonia.

¹⁰ Care should be taken to secure the pure sait Ammonaum saits may contain paradim bases which intrate like ammona but do not reset with besiler a reagent. Fure ammonating sulfate may be prepared by decomposing a high grade ammonatin sail; with sodium hy drouted and p seam, the liberated ammona into pure sulfatine and. The sait is then precipitated by means of sicohol, brought into solution in water and then reprecipitated by alcohol. The final product should be draed in a descence over sulfatine and 17 tidine-free ammonatin salts are now obtainable on the rarket 1 solution containing 0.4714 g of ammonating sulfation in 11 ther of water plus a few drops of conscitutated sulfatine and as preservative contains 1 mg of nitrogen in 10 ml. It is stable indefinitely.

**See 1 preservative*

reading in the colorimeter or photometer. For colorimetric comparisoo, match the standard against itself, and then the unknown against the standard, in the usual way (see p 510). For photometeric measurement, set the photometer to zero density (or 100 per cent transmittance) at 450 mµ (see below) with a blank prepared by diluting 10 ml of Nessler reagent to 100 ml with water This blank will correct for any ammonia present in the diluting water, but not in the digestion reagents, etc For more accurate results, must be a blank and a standard through the entire analytical procedure (digestion, aeration, nesslerization), set the photometer to zero density with the black, and obtain results in terms of the density of the analyzed standard This should correct for ammonia in the reagents as well as systematic errors lother analysis.

CALCULATION For colorimetric measurement

Reading of Standard × mg \ in Standard = mg \ in volume of urine used

At a 1 10 dilution of the unne I ml represents 0 1 ml of urine at a 1.5 dilution 0.2 ml of urine. Yulluply the result of the above calculation by the dilution (10 or a) to obtain the nitrogen content of the unne in mg. per nol or g. per liter (the two are identical). From this the total nitrogeo content of the 24 hour sample may be calculated.

Por photometric measurement

Density of Unknown Density of Standard × mg × in Standard = mg × in volume of urine used

Further calculation is similar to that described for colorimetric measurement

Interpretation. See p 875

Spectrophotometric Characteristics of the Nessler Color with Ammonia. The color reaction between Nessler's reagent and ammonia is the hasis for many other quantitative methods of hiochemical im portance, in addition to its use as described here for the determination of total nitrogen The spectrophotometric characteristics of the Nessler color therefore deserve detailed consideration, since they determine the conditions for accurate photometric measurement, such as the choice of wavelength or filter, solution depth, and range of concentration over which Beer's law is applicable. The data presented here are hased on nesslerization under the conditions described above, i.e. using 10 ml of Nessler reagent in a final volume of 100 ml, in the absence of excess strong acid or alkali or of protective substances such as gum ghatti The values represent the conditions prevailing in the authors' laboratory at the time the analyses were made, and cannot be used for calibration purposes elsewhere, as discussed in Chapter 23, under "Colorimetry and Photometry," hut may be used as a guide for such calibration

The relation between the wavelength at which measurements are made, and light transmittance (optical density) for various amounts of am mona nitrogen, measured at 1 cm solution depth, is shown in Fig 240A. There is no wavelength of peak or maximum light absorption in the visible portion of the spectrum for the amounts of introgen ordinarily encountered and under the presented conditions. The choice of wave-

length therefore depends largely upon the sensitivity desired and the range of concentration over which Beer's law is applicable. This information is supplied by Fig 240B, which represents the data of Fig. 240A, plotted in terms of the relation between transmittance or density and concentration at various wavelengths.

It is shown in Fig 240B that in a 1-em. euvette (or 125-mm. test tuhe, since the two are approximately equivalent) amounts of nitrogen up to 1.5 mg, in 100 ml. of nesslerized solution will read within the accurate portion of the photometer scale (see p. 523) at any wavelength hetween 480 and 520 mμ Such amounts may be determined by calculation based upon the density of a 1 mg standard in accordance with Beer's law, as

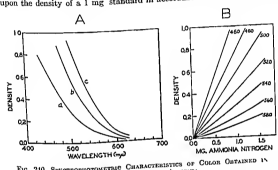


Fig 240. Spectrophotometric Characteristics of Color Obtained in NESSLER REACTION WITH AMMONIA

A, Relation between optical density and wave-length for solutions containing 0.5 mg (a), 1.0 mg (b), and 1.5 mg, (c), of armonia nitrogen per 100 ml final 0.5 mg (c), the control density and 1.5 mg, (c), the control density and volume, at a solution depth of 1 cm. B, Relation between optical density and concentration at the wavelength indicated for each curve

evidenced by the hnearity of the "cahbration curves" over this range. Above 520 mg the sensitivity decreases and slight deviation from Beer's law is noted at higher concentrations of nitrogen Below 480 ma, agreement with Beer's law is excellent over the usable portion of the photometer scale, but the sensitivity is so great that only small amounts of uitrogen may be accurately measured In a 2-cm. cuvette (a 22-mm. test tube is approximately equivalent to this) the same considerations apply, but at only balf the concentration range specified for a 1-cm. cuvette Under these conditions, the standard should contain 0 5 mg of nitrogen, and the unknown not over 1 mg of nitrogen per 100 ml. of nesslerized solution.

The proportionality between density and concentration with the wavelength or filter chosen should be tested for a particular photometer over the concentration range expected; significant deviation from Beer's law greater than that expected from the data of Fig. 240 usually indicated that either the photometer or filter is at fault. In this event, results must be based upon the actual calibration data obtained with the instrument, the use of such a "calibration curve" and the precautions which must be associated with its use, are discussed in Chapter 23

Alternate Aeration Procedures with Iltrimetric Estimation. Instead of nesslenzing the aerated ammona for colorimetric estimation, one may aerate it into acid and make the determination by titration Two such procedures are described. They have the advantage over the colorimetric method of permitting the accurate determination of larger amounts of introgen (up to 3 mg or so). A disadvantage is that too vigorous aeration may carry over a small amount of the fixed alkah used to liberate the ammona, and thus lead to titrimetric but not colorimetric; error. In earth amy spray, the bulb portion of the aeration outlet tube may be loosely packed with these wood, in any event, the apparatus should be tested for this possibility of error by running a blank determination or analyzing a solution containing a known amount of nitrogen.

Aeration into Boric Acid Place 25 ml of 2 per cent boric acid solution containing betweenessed green indicator, as described on p 552 in connection with the determination of blood ures nitrogen in the receiving flask or test tube Liberate the ammonia from the digest by alkali as described for the Folin Farmer method, and aerate the liberated ammonia into the boric acid solution. When aeration is complete remove the receiver and titrate the contents with 0.0143 N sulfuric acid (see p 552) until the more or less blue color is exactly restored to its original yellow green shade, as evidenced by matching against a control 25 ml portion of boric acid plus indicator which is diluted with water to approximately the same final volume as the titrated aample. The end point should be sensitive to about 0.02 ml. of the standard acid.

CALCULATION Lach ral of 0 0143 \ acid is equivalent to 0.2 mg of nitrogen Multi ply the numler of milliliters of 0 0143 \ acid used by 0.2 to give milligrams of nitrogen in the volume of urne user.

Aeration into Standard Acid Aerate the liberated ammonia into a receiver containing 25 ml of 0 01 N sulfuric acid. When aeration is complete, remove the receiver add a drop of methyl red indicator solution, and titrate with 0 01 N sadium hydroxide to an orange or beginning yellow shade

CALCULATION Subtract the number of m of 001 7 alkalu used from 20 0 to obtain the amount of 001 N acid neutralized by the ammonia Multiply this value by 014 to obtain the nitrogen content in mg of the volume of undiluted unne used in the analysis since each ml of 001 N acid repesents 014 mg of nitrogen

3 Determination of Nitrogen by Micro Kjeldahl and Distillation Instead of aerating over the ammonia of a micro-Kjeldahl determination it may be dist lied over and collected in acid for subsequent determination by colorimetric or titimetric methods. In general more accurate and consistent results are obtained by distillation than by acration. Bock and Benediet have described a simple distillation procedures using an ordinary small Liebig condenser? For research work steam distillation has largely replaced simple distillation and is the method of close for all accurate micro-Kjeldahl analyses. Modern stam-distillation apparation (see [1]; 241] is compact.

¹¹ Bock and Benedict J Biol Chem. 20 47 (1915)

casy to use, almost automatic in operation, and requires but a few minutes for each sample

Procedure Digest the sample as described for the Folin-Farmer method, p 876, or digest it with 1 ml of concentrated sulfuric acid plus a few small grains of metallic selenium or a "Hengar granule" (a quartz chip coated with selenium) 2 Digestion must be continued until all of the nitrogen has been

converted to ammonia, the time required for complete conversion varies with different types of material and should be established by trial

Transfer the digested sample with rinsings to the chamber of the steamdistillation apparatus, which has previously been cleared of any contaminating ammonia by a hlank distillation Place the receiving fluid (this varies with the procedure, see below) in a small flask arranged so that the tip of the condenser outlet dips below the surface of the receiving fluid Add sufficient concentrated sodium hydroxide to the digest in the chamber to more than neutralize the amount of acid present (determine by trial) Start the generation of steam in the boiler, and steamdistil the sample until 8 to 10 ml or more of distillate have been collected in the receiving flash Remove the receiver. rinsing down the sides of the condenser outlet tube with a little water in the process, and determine the ammonia in the distribute by any of the following procedures 23

(a) COLORIMETRIC OR PROTOMETRIC ESTI MATION Use 1 to 2 ml of dilute (0 1 N) acid in the receiving flash and collect the distillate as described Transfer the distillate to a container graduated at 25 ml, with rinsings up to about 20 ml volume Add 25 ml of Nessler solution, dilute to 25 ml with water, and mlx

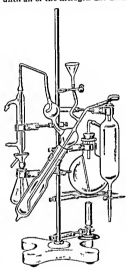


FIG 241 STEAM DISTILLATION AP

Compare in a colorimeter or photometer with a standard containing 0.2 mg of nutrogen, prepared in a second 25-mi container in the same way as for the distillate. In photometric measurement, if the standard is digested and distilled as for the sample, and the photometer is set to zero density with a digested and distilled blank, correction is automatically obtained for any systematic error or for ammonia in the reagents

³² Obtainable from the Hengar Co. Phalodelphia or from dealers in laborator, supj lices. On disconnection or diversion of the steam flow from the clamber of the apparatus the silve of the sample will surelly be removed automatically 13 spil oming in a few morents as the chamber cools down. The apparatus is then really for the next sample without further risangle.

CALCULATION Sim lar to that given for the Folin Farmer procedure p 876 The result gives the mg of nitrogen present in the portion of sample analyzed The conditions for photometric measurement have been presented on p 8.8 For accurate measurement not more than 0.5 mg of nitrogen may be present if a 1-cm cuvette is used or half this amount with a 2-cm cuvette Vuch smaller amounts than this (down to 0.01 mg or less) may be accurately estimated by using 1 ml of receiving and in a test tube graduated at 10 ml collecting the distillate up to the 10-ml mark, adding 1 ml of \csisler solution and comparing photometrically with a suitable standard

(b) TITEIMETRIC LISTIMATION Place 10 ml of 2 per cent bortc acid containing bromocresol green indicator as described on p 552 for the determination of blood urea nitrogen, in the receiving flask collect 10 to 15 ml of distillate as described, and titrate the contents of the receiving flask with 0.0143 V (or 0.02 N) suffuric acid until the solution has been brought back to its original yellow green color Match against a 10 ml control portion of boric acid indicator diluted with water to approximately the final volume of the titrated sample

CALCLATION Multiply the number of ml of 0.0143 N (or 0.02 N) acid used by 0.2 (or 0.23) to obta n the mg, of nitrogen in the portion of sample used Titration of a digested and distilled standard and blank will serve as a basis for correction for any ammonia present in the reagents. Amounts of nitrogen from 0.05 to 3 mg or more may be accurately determined by this procedure. Alternatively the distillate may be collected in 25 ml of 0.01 N and and the excess acid back titrated with 0.01 N alkali in the presence of methyl red indicator as described under the Folin Farmer procedure. The borns and method is preferred because it is a direct titration and requires only one standard solution which is quite stable.

4 Other Methods Many other modifications of the macro-Kjeldahl and micro-Kjeldahl determination of total introgen have been proposed those described here are believed to be as satisfactory as any For micro-Kjeldahl analyses direct nesslerization of the diluted digest has been proposed by Folin and Denis and by Koch and McMeekin these procedures are similar in details to the direct nesslerization methods for the determination of blood nonprotein nitrogen as described in Chapter 23 In general, direct nesslerization is sless satisfactory for unnary mirogen than for blood nonprotein nitrogen because of the difficulty of a voiding turbidity Other methods of determining mirogen include those based upon gasometric estimation their use with unner is similar to that for blood nonprotein nitrogen and references may be found in this connection in Chapter 23

UREA

1 DREASE METHODS

These methods depend upon the principle that the enzyme urease is able at ordinary temperatures to transform urea quiekly and completely into ammonium carbonate Takeuch in 1909 discovered the presence of this enzyme in the soja or soj bean. The application of this enzyme to the determination of urea in urine blood etc., was first proposed by Marshall whose methods have been modified by Van Slyke and Cullen. These latter investigators prepared a permanent preparation of the enzyme, in a water soluble form the use of which makes more con-

venient the rapid and accurate determination of urea in urine, blood and other biological fluids

The urease method is probably the most satisfactory of all methods for the determination of urea. Other nitrogenous constituents such as allan tom are not decomposed by urease. The method involves no carefully regulated heating procedures, and is applicable to diabetic urines. Since however, the basis for the urea determination is essentially an analysis for the ammonia produced by the action of the enzyme, every analysis will include the preformed ammonia which is also present in the urine, so that uncorrected results represent urea plus ammonia To obtain the urea content, it is necessary to know the amount of preformed ammonia present, as established by separate analysis, and to subtract this from the results of the urea determination Urea (and ammonia) determinations should be carried out on fresh or recently collected and preserved samples of urine since ou long standing even in the presence of preservatives significant amounts of urea may undergo bydrolysis to form ammonia thus leading to misinterpretation of the significance of both the urea and the ammonia values. Kinetic studies have shown that a high concentra tion of urea or ammonium ion tends to inhibit the action of urease 24 The conversion of urea to ammonia may take place quite rapidly in urine con taminated with bacteria

a Method of Van Slyke and Cullen Principle. The urine sample is treated with urease and the namionis formed is serated into 0.02 N acid, which is then beek titrated with 0.02 N aliah. A modification based on neration into boric acid and direct titration with standard acid is also described this is preferred over the original method because just as accurate results are obtained and only one standard solution is required in a direct titration.

Parparation of Solid Urease ** Digest 1 part of jack bean meal** with 5 parts of water at room temperature with occas and stirring for an hour and clear the solution by filtration through paper pulp or centrifuging. Pour this extract slowly with stirring into at least 10 volumes of acctone. The acctone dehydrates the enzyme preparation Filter dry in vacuum and pooder. The activity of the preparation is retained indefinitely. Thus prepared it is not perfectly soluble in water, but this fact interferes in no way with its use.

STANARDIZATION OF THE ENZIME PREPARATION Make up accurately a 3 per cent solution of pure uren Treat this solution exactly as the urine is treated in the following method using 0.5 ml of the solution. The armonia formed should neutralize 25 ml of 0.02 N acid or an equivalent amount of acid of other strength. If it does so the preparation is of suffic ent strength to use as indicated. If not more of the preparation must be used for a determination

preveot frothing), and 1 ml of enzyme solution ** Close the tube with a two hole rubber stopper fitted with a long Inlet tube and a short outlet tube, as illustrated in the figures cited, and let the tube stand 15 minutes for the enzyme to act Measure into a second similar tube 25 ml of 0 02 N liCl or 112SO4 Add i drop of caprylic alcohol and 1 drop of methyl red indicator solution Connect the two tubes for aeration with washed air by either pres sure or suction, as described on p 876 At the end of 15 mioutes aspirate for about one half minute to transfer any ammonla present in the free condition to the receiving solution After this aspiration, open the tube containing the sample and introduce 5 ml of saturated potassium carbonate Close the tube at once and aspirate until all the ammonia has been carried over into the acid in the receiver. The time needed for the aspiration varies for different pumps from 5 to 30 mioutes, and should be determined by trial for the par ticular apparatus used At the end of the time needed for the aeratioo, the pump is disconnected (care being taken to avoid back suction) and the excess acid in the receiver is titrated by means of 0 02 N NaOII

CALCLEATION Subtract the number of ml of 002 \ alkali required for the titration from 2.0 to obtain the volume of 002 \ acid equivalent to the aimmonia present Since I ml of 002 \ acid sequivalent to 0.28 mg, of 1002 \ acid equivalent to the aimmonia present is used in an analysis (5 ml of a 1 l0 dilution) multiply the volume of 0.02 \ acid found e pud to the aimmonia by 0.06 (= 2 \times 0.28 to 0.28) to obtain the urea plus aimmonia nitrogen content in g per liter (mg per ml) Subtract from this value the aimmonia nitrogen content in g per liter as established by a separate analysis (see p. 888) to obtain the urea nitrogen content of the urne in g per liter From this and the total volume of the sample the urea nitrogen content of the 24-hour sample is readily obtained.

Agration into Botic Acid Proceed exactly as described above, but agrate the ammonia into 25 ml of the 2 per cent botic acid indicator solution de scribed on p 552 in connection with the determination of blood urea nitro gen (see also the modified Folin Farmer method, p 886). When agration is complete, titrate the botic acid solution with 0 0133. \(\to \to 0.2 \to \to \to \to 1.0

CALALLATION Lach ml of 00143 N and is equivalent to 0.2 mg introgen if 0.02 N and is equal to set of me in ml of and required for it citization 1 y 0.2 (or 0.28) and then further by 2 to obtain the urea plus amonia introgen content of the urine in g per liter Solitzet the ammonia nitrogen content of the urine determined separately to of tain the urina-introgen contint.

Interpretation The mean average daily excretion of urea by normal adults is usually placed at about 20 to 30 g (10 to 10 g of urea introgen) but is very closely dependent upon the protein ingestion and includelism and hence may vary widely in disorders associated with increased tissue catabolism as in fevers the excretion of urea is increased. It may be decreased in pronounced killing and live disorders due to decreased formation but these findings are not

constant The determination of urinary urea, in conjunction with the determination of blood urea, is, however, of major chinical value as an index of ladney function (see urea clearance test. p 965)

The percentage of the total introgen of the unne occurring as urea varies on the average from 80 to 90 On a high-protein diet it is nearer 90 per cent, on a very low-introgen but high-calorie diet it may not be over 60 per cent In marked acidosis it may be considerably decreased relative to the total nitrogen (see "Ammonia," p \$88)

b. Direct Nesslerization Method (Folin and Youngburg¹¹ Modified)
Principle The diluted urine is treated with an alcohole urease solution to convert
urea into ammonia, and the ammonia present is then determined by direct nessleriza
tion. In the original procedure, preformed aumonia is removed by treatment with
Permutit (a synthetic "evchange silicate," see p. 890) and the determination carried
out on the ammonia free solution. It has been found, however, that Permutit may at
times remove a significant amount of the urea as well, and its use has been dispensed
with in the modification described here. The total urea plus ammonia nitrogen is
determined by direct nesslerization after urease treatment, the preformed ammonia
content of the urine, as determined by a separate analysis (p. 888) is subtracted from
the result to give the urea introgen content of the urine.

Direct nesslerization is perhaps not quite so accurate as the aeration procedures because of the possibility of turbidity produced by interfering substances in the color reaction, but the procedure has the advantage of simplicity and the results are quite suitable for most purposes

Procedure Dilute 5 ml of urine to 100 ml. in a volumetric flask and mix well Transfer 1 ml of the diluted urine to a test tube, add 1 ml of the alcoholic urease solution, "and 1 drop of buffer solution "Digest in a beaker of warm water (40° to 55° C.) for 5 minutes or at room temperature for 15 minutes, in the end of which time transfer the contents of the test tube, with rinsings, to a 100 ml volumetric flask, diluting to a volume of about 80 ml Prepare a standard in a second flask, by adding sufficient standard nitrogen solution to contain 0.5 mg of nitrogen," I not of urease solution, and water to a robusto of about 80 ml Trearch flask and 1 ml of geom genetic solution, as et the flask contents to swirling and add from a graduated cylinder 10 ml of Nessler reagent "Dilute immediately to 100 ml mlx, and allow to stand 5 minutes Read in a colorimeter or photometer in the usual way For photometric measurement, set the photometer to zero density with a blank prepared similarly to the standard except that the standard nitrogen solution is amitted.

CALCULATION For colorimetric measurement

 $\frac{\text{Reading of Standard}}{\text{Reading of Unknown}} \times \frac{\text{mg N in}}{\text{Standard}} \times \frac{\text{Dilution}}{\text{Dilution}} = \frac{\text{mg}}{\text{m I ml}} \text{ of undiluted urine}$

³⁴ Folin and Youngburg J Biol Chem. 38 111 (1913) and Youngburg J Biol Chem. 45, 391 (1931).

ii To prepare the alcoholic urease solution place 3 g of Permutt in a final, wash once with 2 per cent acetic acid then twice with water add 5 g of fine jack bean med and 100 ml of 30 per cent alcohol Shake gailly but continuously for 10 to 15 minutes and filter The filtrate contains practically all of the urease and extremely little of other materials in Dissolve 14 2 g of Naill 10, and 12 g of Naill 10, and 12 g of Naill 10.

[&]quot;Dissolve 142 g of Naill O. and 120 g of Naill PO. or equivalent amounts of the crystalline salts in water and make up to 100 ml

[&]quot; See footnote 19 p 977

[&]quot; See Appendix

Subtract from this value the ammonia content of the urine, as determined separately, to obtain the urea nitrogen content of the urine. Under the conditions described, the dilution is 20, and urines containing 5 to 20 g of urea nitrogen per liter may be accurately analyzed. For values outside this range, repeat the analysis at n more satisfactory dilution.

For photometric measurement

Density of Unknown Density of Standard X Dilution = mg urea- plus ammonia nitrogen in 1 ml undiluted urine

Correct for the preformed ammonia content as described above. The conditions for photometric measurement of the Nessler color have already been described (p. 878) At 480 m_H, and in a 1-cm cuvette, the standard has a density of approximately 0 400, permitting measurement with unnes containing up to about 25 g of urea nitrogen per liter under these conditions. For higher values, or for photometric measurement at greater depth of solution, use a greater debt dultion of the urine

Interpretation. Sec p 884

c. Microdiffusion Method of Conway." Principle. The sample of diluted urine is incubated with urease in the outer compartment of a "Conway



diffusion cell" (Fig 242) "The ammonia formed is then liberated by addition of saturated potassium carbonate solution, and the cell is covered The ammonia diffuses over into and is absorbed by acid in the inner compartment of the cell, where it may there be determined by itimmetric or colorimetric means. As with all urease methods for urine, separate determination of the ammonia N present is necessary in order to obtain the urea N content.



Fig 242 Conway

The same principle may be employed for the determination of urea N in blood, of total N in micro-Kieldahl digests, and in general wherever a volatile absorbable substance is to be determined (see p 668)

Because of the simplicity of the procedure and its application to microanalysis, increasing use is being made of this principle in routine and research laboratories

Procedure. Determination of Urea N Plus Ammonia N. Dllute 1 ml. of urine to 10 ml. with water and mir. With an accurate pipet transfer 0 2 ml. of this diluted urine to the outer compartment of a Conway diffusion cell (Fig. 242). Place 1 ml ol 2 per cent boric acid solution containing added bromocreaol green indicator. In the inner compartment S mear a glass cover plate with 2

¹² Conway and Byrne Biochem J 27, 419 (1933) Conway biol. 27, 430 (1933) The procedure as described here is a slight modification of the original Conway procedure and includes somie suggestions of 'tenuts U Lob Clin Med 23, 228 (1933-1919)) For application of the microdiffusion principle to the determination of annial amounts of application of the microdiffusion principle to the determination of annial amounts of the conversation of the determination of the dete

[&]quot;The unit shown here is made of Coors porcelain and may be obtained from the Arthur H. Thomas Co Philadelphia.

is See p. 552 for preparation of this solution

thin film of fixative is and place on the cell, greased side down. Slide the cover aside far enough to permit the introduction of a pipet tip, and add to the outer chamber 0.5 ml, of urease-phosphate solution. 37 Replace the cover, and mix the fluids in the outer chamber by slight tilting and rotation of the cell. Set aside at room temperature for 15 minutes. At the end of this time, tilt the cell slightly to displace the fluid in the outer chamber to one side, slide the cover aside slightly to permit the introduction of a pipet at a point opposite to the displaced fluid, and add 1 ml. of saturated potassium carbonate solution to the outer chamber. At once replace the cover, mix the fluids in the outer chamber by tilting and rotation as above, and set aside for 90 mlnutes at room temperature (or place in an incubator at 38° C. for I hour). Be sure the cover is firmly in place during this period. At the end of the period, remove the cover and titrate the contents of the inner champer with 0.0143 N sulfuric acid. s using a microhuret with a fine tip which dips below the surface of the solution being titrated. Titrate until the color of the fluid In the inner chamber exactly matches that in a control celi. The control consists of a second cell treated exactly as described for an analysis except that 0.5 ml, of water replaces the urine sample.

Ammonia N Determination. Proceed as ahove, hut use 0.2 ml. of undiluted urine and omit the treatment with urease, adding the 1 ml. of saturated potassium carhonate solution directly to the urine in the outer compartment, allowing to stand covered for diffusion and absorption of ammonia to take place, titration, etc., as described.

CALCULATION, Since 1 ml of 0 0143 N sulfure acid is equivalent to 0.2 mg, of oitrogen, the calculation is as follows

- a. Urea N + Ammonia N.
- mi 0 0143 N acid required × 0 2 × 50 = mg urea N + NH, N per ml

undiluted urine

- b Ammonia N.
- ml 0 0143 N acid required \times 0 2 \times 5 = mg NH₂ N per ml undiluted urine
 - c Urea N subtract (b) from (a) to obtain the urea N content of the urine, in mg. per ml (or g. per liter)

During the titration, the fluid to the moer compartment should be stirred continuously, a convenient arrangement consists of a fine glass tip delivering compressed air to a slow stream of bubbles below the surface of the solution. A oragnetic stirrer acting on a "flea" (small piece of 100 were enclosed in a small sealed glass tube) placed in the inner compartment is also an excellent method of stirring during the titration. If desired, the determination may be made colorimetrically or photometrically by placing 1 ml. of 0 1 N sulfurn end (without nodeator) in the more compartment,

⁴⁴ Ordinary petrolatum is satisfactory at room temperatures. If the cells are to be incurred at 35° C, a petrolatum-paraffin wax maxture is recommended. Melt 3 parts petrolatum with 1 part paraffin wax (MP 55°) and allow to cool.

¹⁷ Dissolve 400 mg of urease powder (prepared as described on p 883, commercial preparations may also be used) in 10 ml of water and add 0.4 ml of phosphate buffer (0.9 g of Naili-PO, II-O and 17.9 g of NailI-PO, 12II-O in 100 ml of water) Prepare the urease solution fresh daily, and discard any unused portion. The activity of the powder may be tested by a control analysis on pure urea solution. Dilute a 2.14 per cent solution of urea 1:10 with water and analyze 0.2 ml, of the diluted solution, 0.2 mg, of nitrogen should be found.

³ Dilute 14 3 ml, of N sulfuric acid to I liter in a volumetric flask and mix. This solution is quite stable.

instead of the boric acid solution. After absorption of ammonia is complete, the con tents of the inner chamber are transferred by means of a rubber bulb pipet, with rusings, to a test tube graduated at 20 ml and nesslerized at this volume in the presence of 2 a ml of Nessler reagent Colonmetric or photometric estimation may then be made in terms of a standard containing 0.2 mg of nitrogen, nesslerized similarly

Interpretation. See previous methods

2 OTHER METHODS

Many other methods for the determination of urmary urea content have been described some of which combine certain features of the vari ous procedures described here Rose and Coleman suggested colonmetric determination of the acrated ammonia (as in the Folin Farmer micro-Kjeldahl metbod, p 876), after treatment of the urine with urease Marshall described a simple chincal method based upon direct titration of the ammonia liberated by urease treatment 29 Gasometric measurement in the Van Slyke-Neill apparatus (p 709) of the carbon dioxide liberated by the action of urcase on urea40 is simple, rapid, and accurate, the only disad vantage is when many analyses are to be done, since each analysis must be carried to completion before the next is begun. Gasometric determination hased on the liberation of nitrogen by treatment with hypobromite has been described by Stchle 41 Methods based upon isolation of urea as the insoluble xanthydrol compound have been described by Barrett and Jones and by Allen and Luck "Direct colorimetric methods which do not involve the action of urease have been proposed by Ormsby," Barker, 45 and Archibald 46

AMMONIA

Introduction. Many of the methods which have been proposed for the determination of urmary ammonia are based upon principles similar to those already described in connection with the micro-Kieldahl deter mination, and the determination of urmary urea since these const t essentially in the determination of aminonia produced as a result of the analytical treatment. For the accurate determination of urinary ammonia it is necessary to separate the ammonia from interfering material, and to avoid the use of excessive heat or strong alkali in the treatment of the urme, since these may lead to the production of ammonia from other nitrogenous constituents of the urine. The successful separation of am monia from urine by aeration was first described by Folin, 47 his method was modified and improved by Van Slyke and Cullen and their method (with modification) is described here since it requires less urine than Folm's method, gives just as accurate results and utilizes the same appa-

³¹ Marshall J Biol Chem 14, 283 (1313)

[&]quot; Yan Slyke J Buol Chem 73 603 (1927)
" Stelle J Buol Chem 47 13 (1321) and 51 83 (1322) See also Yan Slyke J Buol. Chem \$1 449 (1929)

Harrett and Jones Biochem. J 24 1246 (1J32)
 Allen and Luck J Biol Chem. \$2 (93 (1J29)

[&]quot;Ormsly J Biol Chem 144 405 (1912 "Barker J Biol Chem. 152, 453 (1544)

⁴ Archibald J Biol Chem. 157 407 (1315) of Folin's method is described in the Fleventh Edition of this book.

ratus, reagents and technique as have already been described for the determination of urea in blood (p 551) and in urine (p 883)

1. Aeration Method (Van Sij ke and Cullen, Modified) Principle The urine is treated with an equal volume of saturated potassium carbonate solution, and the liberated ammona is transferred by aeration into an ear receiving solution, where it is then determined by ittration. In the original procedure, the ammonia is aerated into 0.02 N acid which is then back titrated with 0.02 N alkali. In the modification described here, boric acid solution is used to receive the ammonia, which is then titrated directly with standard acid. Results are the same by either procedure, the boric acid method has the advantage of being a direct titration and requires only one standard solution which is quite stable.

Procedure Measure 5 ml of undiluted urine into one of the two large test tubes used in an aeration train (see Fig. 238, p. 877, and Fig. 147, p. 552) and connect this tube for aeration, as shown in the illustrations, with a second tube containing 25 mi of the 2 per cent boric acid containing added bromocresol green indicator, described on p 552 in connection with the determination of blood urea nitrogen. Add a drop of caprylic alcohol to each tube to minimize foaming When ready, remove the stopper of the tube containing the urine and add 5 ml of saturated potassium carbonate solution. Replace the stopper tightly and start the air current (pressure or suction, the incoming air must be washed by preliminary passage through a wash bottle containing dijute (1 10) sulfuric acid, to remove any ammonia present) The air current should be run slowly for the first two minutes, and then increased to a rate as fast as the apparatus will stand Aeration is continued until all the ammonia has been driven over, this may take from 5 to 30 minutes depending upon the apparatus, and the time required should be established by trial

When aeration is complete, remove the tube containing the boric acid, rinsing down the inlet tube in the process, and titrate the contents with 0 0143 N (or 0 02 N) sulfuric acid, until the more or less blue color is replaced by the original yellow green color, as determined by matching against a control 25 ml portion of the boric acid indicator solution which has been diluted with water to approximately the final volume of the titrated sample. The end point should be sharp to about 0 02 ml of standard acid

CALCULATION Each ml of 0 0143 N need is equivalent to 0.2 mg of ammonia nitrogen, if 0.22 N acid is used, 1 ml equals 0.28 mg of ammonia nitrogen. Multiply the volume in ml of acid required for the tirration by 0.2 (or 0.28), to obtain the ammonia-nitrogen content of 5 ml of urine divide the result by 5 to obtain the ammonia mitrogen content of the urine m mg per ml (or g per liter) Results on any other basis (e.g., g. per 100 ml or per 24-hour sample) may then be readily obtained

If the original Van Sjyke-Cullen procedure based on back utration of 0 02 N and with 0 02 N alkali is used the conditions and the calculations are similar to those described on p SS3 in connection with the determination of urmary urea except that results are divided by 10 because 10 times as much urms is used for the aminoma determination as for the urea determination.

The ammonia acrated into dilute (902 Å) sulfaire and may be nex-lettred and determined colorimetrically or photometrically as described for the Folia Farmer micro-hyeldahl procedure p 876 this procedure was suggested by Folia and Mac-Cillum The amount of urine taken should not contain over 0.5 to 10 mg of ammonia nutrogen; i.e. 1 to 2 ml of normal urine or less in acidosis diluted to about o nul, with water Comparison is made against a suitable standard ness-lettred in the same way, and the calculations are similar to those for the Folia Farmer procedure

Interpretation. The average daily output of ammonia nitrogen in the urine of an adult on a mixed diet is about 0 7 g, corresponding to about 50 milliequivalents, or 500 ml. of 0.1 N base, per day, and representing about 2.5 to 45 per cent of the total nitrogen. The amount excreted may vary widely from this average value, however, since (along with the titratable acidity of the urine, see p. 869) the ammonia content of the urine appears to parallel the state of acid-base balance within the body. It is increased in amount by the ingestion of acids or acid-forming foods and in the acidosis of starvation or diabetes (but not of nephritis), and decreased by the ingestion of alkalies or base-forming foods and in alkalosis. The ammonia of the urine is liberated enzymatically in the kidneys principally from glutamine but also from amino acids and other possible precursors Urmary ammonia appears to function as a "synthetic" base, capable of replacing "fixed base" (sodium, potassium) in the excretion of acids as their neutralized salts, and thus conserving such fixed base to the organism.

2. Formal Titration Method (Malfatti): Principle, This method is based on the reaction that takes place when formalin solution is added to a solution containing ammonium static (see "Amino Acid Nitrogen" methods). An acid reaction is produced in the mixture, which is then ittrated with standard alkali using phenolphthalein as an indicator. Animo acids give the same reaction so that the result of the titration represents ammonia + amino acid nitrogen. This method may be used for the rapid clinical estimation of these forms of nitrogen as a substitute for an ammonia determination, but the results do not represent ammonia as is smeltimes stated.

Procedure. To 15 ml. of urine in a 200-ml. Erlenmeyer flask add 5 g. of finely pulverized potassium oxalate, a few drops of phenolphthalein, and turste to a faint hut permanent pink color with 0.1 N NaOH. (The urine muture just after neutralization in the urinary titratable acidity determination may he used.) (See p. 870.) Then add 10 ml. of neutral formain solution (see "Amino Acid Nitrogen"), mix well, and titrate with 0.1 N sodium hydroxide to a permanent pink color.

CALCLIATION One ml. of 01 N sodium hydroxide is equivalent to 17 mg. of ammonia. Multiply the number of ml of 01 N alkali required after the addition of the formalin hy 17 and by 4 to get the number of mg of ammonia + amino acid nitrogen

(expressed as ammonia) in 100 ml of the urine examined

3. Permutit Method: Principle. The urine is shaken with particles of an "exchange sheate," which remove the ammonia from solution. The ammonia is set free from the sheate by treating with alkali solution. This is then nesslerized and compared with a standard ammonia solution nesslerized in the same way.

Procedure. Introduce about 2 2. of Permutit powder into a 100-ml. volumetric flask. Add about 5 ml. of water (no more), followed by 2 ml. of urine, "

⁴ Folin and Bell J Biol Chem. 27, 329 (1917) Permutit is a synthetic aluminum sheate which has the property of taking up ammonia quantitatively in neutral or weakly acid solution and of releasing it in strongly althain solution. The 60-80 mesh powder about the strong of the folial solution. The 60-80 mesh powder about settle quietly leaving the supernation fluid early active acid, and finally with water gazantee used at the washed with water, 2 set early acid and finally with water gazantee.

[&]quot;With urmes very low in ammonia it may be necessary to use more urine (5 ml) but so far as it is practicable it is better not to use more than 2 ml and to employ a weaker

accurately measured. Rinse down the added urine by means of a little water (1 to 5 ml.), and shake gently but continuously for 5 minutes. Rinse the powder to the bottom of the flask by the addition of water (25 to 40 ml.) and decant. Add water once more and decant. (In the case of urines rich in bile it is advisable to wash once or twice more) Add a little water to the powder, introduce 2 ml. of 10 per cent sodium hydroxide, shake for a few moments and set aslde, while preparing the standard ammonium sulfate solution as follows.

Transfer 10 ml. of the standard ammonlum sulfate solution (see p. 877) containing 1 mg, of nitrogen for 5 ml, containing 0.5 mg, of nitrogen if a half-strength standard is desired) to another 100-mi volumetric flash and add 2 ml of 10 per cent sodium hydroxide (to balance the aikail added to the Permutit mixture in the other flask), Dilute to about 75 mi, and mix, Transfer 10 ml, of Nessler's solution (see p. 877) to a measuring cylinder. Now give the volumetric flask a vicorous whirl so as to set the solution spinning within the flask and add at once the whole of the Nessler solution in the cylinder. With another whiring movement complete the mixing of the contents of the flash, if the process of nesslerization has been successful a deep-red but crystal-clear solution is obtained, If it is not perfectly clear discard it and prepare a fresh standard. Then dilute the contents of the flask containing the Permutit and the urmary ammonia to about 75 ml., whirl the mixture, and add the Nessler reagent (10 ml) exactly as in the case of the standard solution, Dilute the contents of both flasks to volume (100 ml.) and compare in a colorimeter with the standard set at 20 mm, or determine the densities of unknown and standard in a photometer at 480 mu For photometric measurement, set the photometer to zero density with a blank prepared by treating about 75 ml. of water plus 2 ml of 10 per cent sodium hydroxide in a 100-ml. volumetric flask with 10 ml. of Nessler solution as described, diluting to 100 ml , and mixing.

CALCULATION For colorimetric measurement

Reading of Standard \times mg N in Standard \approx mg NH, N in volume of urine used

Divide the result of the above calculation by the volume of urine used to obtain the ammonia-nitrogen content of the urine, in mg per ml (or g per liter) From this, and the volume of the 24 hour specimen, the daily output of urinary ammonia is readily obtained.

For photometric measurement

Density of Unknown X mg N in Standard ~ mg NH₂ N in volume of urine used

Further calculation is the same as for colorimetric measurement. The spectrophotometric characteristics of the Nessler color, and the conditions for accurate photometric measurement, are similar to those given on p 578 in connection with the Folin-Farmer nucro-Kicklahl method, since similar amounts of nitrogen are dealt with here, under convision to orditions.

Interpretation, See p 890

standard (0.5 mg instead of 1 mg of mitrogen) for colorimetric comparison. If there is doubt about which standard to use at its sound analytical practice to prepare both and to match the unknown against the most suitable standard as determined by inspection

AMINO ACID NITROGEN

1. Ninhydrin Method of Von Slyke, MocFodyen, and Homilton:* Principle. The urne sample, previously freed from urea by treatment with urease, is heated at 100°C in a closed reaction vessel with minhydrin (see p 129). Ammo acids present yield carbon dioxide quantitatively under these conditions. The carbon dioxide produced is transferred to the chamber of the Van Slyke-Neill manometric apparatus and there determined From the amount of carbon dioxide found, the e-amino nitrogen content of the sample is readily obtained, since all of the common amino acidy god or-amino introgen present. Proteins, peptones, peptides (other than glutathiose, which has a free e-amino group), and substances other than amino acids do not read significantly. This procedure is considered to be the most specific yet devised for free amino acids.

Procedure: Removal of Urea and Preformed CO₂, Place 2 mi. of fresh urine (or urine preserved by saturation with thymoi and storage in the cold) in the all-glass reaction vesself (Fig. 233) and add if orgo of 0.04 per cont bromothymol blue. if it a blue color is obtained, add 1 N sulfuric acid drop by drop until color turns yellow, and then 1 N sodium hydroxide until just blue. If a yellow color is ohtained initially, add 1 N sudmin hydroxide until just blue. Then add 175 mg. of dry phosphate bufler mixture, 10.2 ml of a 1 per cent solution of urease, 1 and a crystal of thymol Stopper loosely to prevent loss of water but not of carbon dioxide and incubate overnight at 3.7 to 40 °C. After incubation add 1 drop of 0 04 per cent bromocresol green and a drop

of caprylle aicohol. Cautiousiy add 5 N sulfuric acid drop by drop, with gentle whirling to minimize foaming, until the solution is just sellow (ni alout 3), then add 100 mg of dry citrate buffer mixture. Made several pieces of alundum to prevent bumping and heat to boiling over a microburner.

<sup>ω Van blyke, MacFadyen, and Hamilton J Bud Chem 150, 251 (1943) See also
Hamilton and Van blyke J Bud Chem 150, 231 (1943) Van Slyke Dillon MacFadyen
and Hamilton tod. 141, 027 (1941) Van Slyke and Folch tod. 156, 509 (1940)</sup>

¹¹ Obtainable from E. Machlett and Son New York N Y
11 See Chapter 1 for the preparation of indicator solutions

ii Dry Phosphate Buffer Misture Grind separately in a mortar 3 parts by weight of anhydrous incompostassium phosphate and 1 part of anhydrous disodium phosphate (or 25 parts of theodium phosphate containing 12 molecules of water). Mix the two ground solide and grind intimately together Dispense with a glass spoon calibrated to deliver 175 ± 15

ing of the powder This mixture produces a GI of 6 2 when in solution

1. Urease Solution This is made from urease powder prepared as described on p. 583.

Dissolve the powder in water in the processor powder prepared as described out has 10 per cent solution 1 10 with water before use to make the 10 ml of water before use the prepared to 10 ml of water before use the produce a correction (as established by the blank analy as described in the procedure) equal to several per cent of the usual anno and one tent of the unal anno and content of the unal anno and the unal anno and the unal anno and the unal anno and the unal anno and the unal anno and the unal anno and the unal anno and the unal anno and the unal anno and the unal anno and the unal anno and the unal anno and the unal anno and the unal the unal the unal the unal the unal the unal the unal the unal the unal anno and the unaliteration of the unal the unal the unal the unal the unal the una

solution than I per cent should be used to use per manute a proportional() strong by the burst Buffer (firster Grand separately in a mortar 206 g of sodium citrals (NacCillO 2110) and 1915 g of citra scal (CillO; 110). Win the two ground solid

an I grin I intimately together Disputs with a glass spoon calibrated to deliver 100 ± 10 mg, of the powder. This nurture produces a pH of 2.5 in solution.

Boll exactly 1 minute (no longer). Cool to below 25° C. (placing in ice water for three minutes is satisfactory). Attach the short rubber connector to the side arm (Fig. 243), and wipe off any water from the inside ground-glass surface of the ressel top.

Treatment with Ninhydrin. Add 100 mg. of ninhydrin¹¹ to the cooled reaction mixture with the aid of a small funnel having a short stem. Have the glass stopper ready with a thin film of special lubricant¹³ spread over the

ground-class surface, and as soon as the ninhydrin has been added, remove the funnel and set the stopper in place, with the hole in line with the side arm, as shown in Fig. 243. Immediately attach the side arm to a good source of vacuum (preferably with attached manometer) and evacuate quickly to 30 mm, or less pressure, shaking gently if necessary to minimize effervescence. When the vessel is evacuated (10 seconds may be sufficient time), turn the stopper through 180° to close off the vessel. Press the rubber connector flat with the fingers, detach vessel and connector from the vacuum line, and insert a glass plug moistened with water or glycerol into the connector, trapping as little air as possible In the process. Tighten the stopper in place-it may be held more securely by a heavy rubber band linking stopper and side arm Place the closed vessel unright in a wire rack and immerse completely in vigorously boiling water for 8 minutes

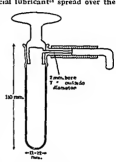


FIG 243 REACTION VESSEL FOR MINHYDRIN ANINO ACID METHOD Courtes: Hamilton and Van Slyke J Biol Chem 150 231 (1943)

Remove, tighten the stopper again by a slight twist if necessary, and allow to cool to room temperature. The closed vessel may be kept for several days at this point if necessary, and if many analyses are to he done they may all be brought to this stage before proceeding.

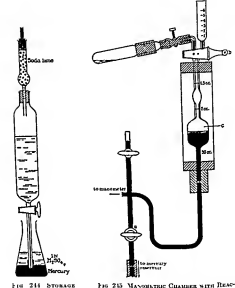
Absorption and Measurement of CO₂. Warm the contents of the reaction vessel to 38° to 48° C, by immersion in a beaker of water at this temperature for 10 minutes While waiting, deliver 2 ml of the carbonate-free 0 5 N sodium hydroxide in sodium chloride solution³ from the storage container

¹⁴ A 5-cm length of stethoscope tubing as used (3 mm bore 12 mm outside diameter) Before using for the first time, clean the tubes maske with a test-tube brush immerse in acidified water in a round bottomed flash, and bod for 30 minutes. Remove the flame, stopper the flask and cool under the tap. The vacuum caused by cooling draws excess gases from the rubber. When hubbles no longer appear, open the flask and wash the tubes with water. One treatment suffices.

ar Dispense with a glass spoon calibrated to deliver approximately the required amount (within 10 per cent)

¹⁴ A lubricant which does not leak at high temperatures must be used. New statum N.Y. heavy lubricating grease obtainable trom E. Machlett and Son New York is satisfactory. A suitable lubricant may be prepared in the laboratory as follows Mix 35 g of alumnum distearate to a paste in 100 ml of heavy parafilm immeral oil. Heat with continuous stirring in a beaker over a low flame to effect solution of the soap Allow to rool then work up the frable gel to a smooth translucent paste on a glass plate with a steel spatule preferably with warrung to 45° to 50° C.

[&]quot; Dissolve solid sodium hy droxide in an equal weight of water and allow to stand until



CONTAINED FOR ALKA-LIVE SOLUTIONS. HUNG WITH THE PRO-TECTED PROM ATMOS-PICERIC CARRON DI-OXIDE

Courtesy lan lyis and licirly J Head Chem., 136,

509 (1940)

TION VESSEL (CARLIER TYPE) ATTACHED BUT NOT YET CONNECTED WITH INTERIOR OF CHAMBLE Courtesy Van Slyke Dillon MacFadyen, and Hamil

ton J Bul Chem 141, 627 (1941)

the carbonate settles Standardize the supernatant solution by pipeting 7 ml. into water and iterating with standard (2 N or stronger) and From the result, determine how much of the I I NaOli is required to make 250 ml of 0.5 N solution. Fill a 250-ml, volumetric flask to within about 10 ml of the mark with concentrated CO-free NaCl solution (dissolve 250 g of VaCl in 750 ml of water deserate and store as described on p. 715). Pipet the correct amount of 1 1 NaOH solution into the flask delivering it below the surface of the salt solution. tild a few drops of I per cent alizarin solution as indicator, fill to the mark with salt solution, and nut. Transfer to a storage container (Fig. 244 in text) to protect from atmospheric carton dioxide

equipped with a rubber tip (Fig. 244), into the chamber of the Van Siyke-Neili apparatus (Fig. 188), through a mercury seal in the cup (see Fig. 190 and p. 711 for method of delivery). Seal the cup capillary with a little mercury, lower the mercury in the chamber to about the middle of the chamber. and close the cock leading to the leveling bulb (cock a in Fig. 245). By this time the reaction-vessel contents should be sufficiently warm. Remove the vessel from the water hath, remove the glass plug from the side arm, attach the rubber connector to a source of vacuum, and reevacuate the side arm for

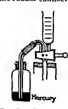


FIG. 246. SMALL BOTTLE OF MERCURY ARRANGEO FOR SEALING CAPILLARY OF MANOMETER STOP-COCK.

Courtesy, Van Slyke and Folch. J. Biol. Chem, 136, 509 (1940).

a few seconds. Pinch the rubber connector flat with the fingers, detach from the vacuum line, and at once attach to the side arm of the Van Sivke-Nelli apparatus. Fig. 245 shows the conditions prevailing at this point. Turn the stopcock of the reaction vessel and cock b of the chamber so as to connect vessel and chamber and permit gas to pass from one to the other. Transfer the CO, in the reaction vessel to the chamber (where It is absorbed by the alkali present) by repeatedly raising and lowering the mercury in the chamber by manipulation of the leveling bulb and cock a. The first lowering of the mercury must be done carefully to prevent bumping in the reaction vessel, and at each lowering the reaction vessel is shaken by hand to distribute the fluid along the wails. Five excursions of the mercury suffice to transfer all the CO, each excursion taking about 10 seconds, After the last upward excursion, lower the mercury in the chamber to about the middle point, and close cocks a and b.

Remove the reaction vessel and fill the side arm of the cock capillary of the manometric apparatus with mercury by suction from a small bottle containing mercury (see Fig. 246).

Open cock a, raise the leveling bulb to about even with cock b, then close a. Open b carefully to the right-hand cup at the top, allow the gases present to escape through the cup, then admit mercury to the chamber through cock a until the fluid in the chamber just reaches the top of the chamber, at cock b. Close a and then b. Place the leveling bulb at its normal position (approximately even with the 50-ml. mark at the bottom of the chamber), place a little mercury in the cup, and seal the cock capillary with mercury in the usuai way.

Through the mercury seal, add exactly 1 ml. of 2 N lactic acid in sodium chioride solutions from a stopcock pipet provided with a rubber ring on the tip, as illustrated in Fig. 190, p. 711, then seal the cock with a little mercury. By manipulation of the leveling bulb and cock a, lower the mercury In the chamber to the 50-ml. mark, close a, and shake the chamber for 20 to 30 seconds. The liberated CO: drives the mercury below the 50-ml. mark Raise the leveling hulb, open a carefully, and adjust the mercury level to the 50-ml, mark, Close a and shake the chamber for 3 minutes. Place the leveling huib high, open a, and admit mercury into the chamber until the level of the fluid (not mercury) meniscus is exactly at the 2-ml. mark on the chamber, This adjustment must be done in a consistent manner. Allow the mercury to enter through a steadily, without jerky oscillation of the chamber contents.

[&]quot;Dilute 1 volume of concentrated factic acid (sp. gr. 1.20) with 4 volumes of the concentrated NaCl solution described in the previous footnote,

and complete the adjustment in 30 to 40 seconds. If the adjustment is missed the first time (or for a duplicate reading) lower the mercury to the 50 ml mark, shake for one minute, and repeat. With the fluid meniscus at the 2-ml mark, read the manometer. This reading is p.

Replace the leveling bulb to the normal position, open cock a, and place 0.5 ml of 5 \ sodium hydroxide solution in the cup above cock b Admit the alkali to the chamber slowly, over a period of about half a minute, until only enough is left to fill the capillary beneath the cup Add a little water and mercury to the cup, and seal the cock with mercury If the viscous alkali forms a solld column in the top of the chamber, it may be broken up by admitting a little mercury in abort bursts. Wix the alkali and ensure completed absorption of CO₁ by raising and lowering the mercury in the chamber 3 times in short excursions, then hring the fluid mensions to just below the 2 ml mark no the chamber Allow I minute for drainage, then adjust exactly to the 2 ml mark. Read the manometer, this reading is p.

CALCULATION The pressure Pcor due to the carboxyl carbon dioxide of the amino acids present is found as follows

$$P_{CO1} = p_1 - p_2 - c$$

where e is the correction due to the blank. To establish e run through an entire analy sis as described above (including incubation with urease etc.) on 2 ml of water in stead of urine. The difference hetween the p₁ and p₂ readings for the water blands analysis equals e. This blank will serve to correct for small amounts of amino acids which may be present in the urease preparation as well as for certain possible variables in the manometric measurement likely.

To obtain the amino acid nitrogen cootent of the sample multiply Pcot by the proper factor, obtained from the accompanying table

Pcox X factor = mg. a-amino nitrogen per liter of urine

The temperature column in the table refers to the temperature of the water in the jacket surrounding the chamber at the time the p₁ and p₁ readings are made and a is the volume of the gas phase when the manometer is read a is ordinarily 2 mil but may be reduced to 0.5 ml u only small amounts of amino acids are present.

FACTORS FOR USE IN CALCULATING MG OF Q-AMINO NITEOGEN FER LITER OF URINE (SAMPLE VOLUME = 2 Mr.)

•		_		
Ten persisse * (—	a = 20	a = 05	Temperature a = 20	a = 05
15 16 17 18 13 20 21 22 23	0 802 0 738 0 735 0 732 0 783 0 786 0 782 0 780 0 776	0 2010 0 2002 0 1994 0 1385 0 1367 0 1369 0 1364 0 1353 0 1915	25 0 7.0 26 0 767 27 0 764 28 0 762 29 0 7.8 30 0 7.6 31 0 -52 32 0 7.0 33 0 7.47	0 1930 0 1923 0 1916 0 1908 0 1901 0 1894 0 1886 0 1873 0 1873
24	0 773	0 1938	31 0 744	0 1866

⁴⁴ Mix 1 volume of t t \a011 solution with 2 volumes of water

Interpretation. The a-aming acid mitrogen content of normal urine appears to make up about 1 per cent (or possibly even less) of the total urmary nitrogen, corresponding to a daily exerction of 100 to 150 mg of α-amino introgen Results by other methods, e.g., the introus acid method of Van Slyke and Kirk, 6° the formol titration procedure (see following method), and the copper titration procedure of Albanese and Irby, 63 tend to he either slightly or significantly higher than results by the junhydrin procedure described here, presumably because of differences in specificity for α-amino nitrogen Little is known concerning the significance of normal or pathological variation in amount of urmary a amino nitrogen Excretion may be largely increased in disorders associated with tissue waste as typhoid, in pronounced atrophy of the liver, acidosis, etc

Microhiological assays for individual free amino acids in human urine show a preponderance of histidine, cystine, and glycine, average daily excretions heing 150 to 250, 60 to 110, and 120 mg, respectively Threomne, 50 to 60 mg, and lesser amounts of other amino acids are also found 64 Glycine, glutamic and aspartic acids, and lysine are also found

in conjugated form

2 Henriques-Sørensen Formol Titration Method 15 Principle A solution containing amino acids is a early neutral in reaction. If formaldehyde be added, how ever, a reaction takes place with the formation of derivatives which are more strongly acid in reaction due to the destruction of the basic properties of the amino groups (see p 130 for discussion and equations) The carboxyl groups may then be titrated using phenolphtbaleia as an indicator. The acidity as shown by the titration is a measure of the amount of amino acid nitrogen present. Ammonia likewise reacts with formalde hyde in a similar manner as is shown in the following equation

$$4NH_4Cl + 6CH_4O = N_4(CH_4)_6 + 6H_4O + 4HCl$$

Hence the formal titration in the presence of ammonia gives results which include both amino acid and ammonia nitrogen. Ammonia may be determined and a correction applied or the aminoma may he removed by means of phosphotungstic acid Phos phates also interfere by obscuring the end point and are removed by the addition of harium salts Polypeptides and still more complex protein derivatives likewise react with formol to a certain degree so that the results do not strictly represent amino acid hitrogea "

The method is, with some modifications involving the preparation of the solution to be titrated, applicable in the determination of amino acids in any medium, eg, urine, protein digests, to etc. The presence of buffers or poorly dissociated acids, eg, some fatty acids, will tend to give values which are too high Certain of the amino acids when present in large amounts will give erroneous results, but in the ordinary urine or digest these errors are either negligible or compensate each other. In the titration of colored solutions the control solution which is necessary in this method must be

⁴¹ Van Sixke and Bark J Biol Chem 102 651 (1933)

[&]quot;Hanese and Irby J Bol Chem 13 53 (1944)
"Dunn Camen Shankman and Block Arch Bochem 13 -07 (1947) Gutman and Dunn Camen Shankman and Block Arch Bochem 13 -07 (1947) Gutman and Lynch and Merander J. Bol Chem 165 527 (1941) Christensen Cooper Johnson and Lynch bid 1941 (1947) Shefiner hissner and I almer bid 175 107 (1948) Shefiner hissner and I almer bid 175 107 (1948) See also Northrop J Gen Ph. Henriques and Sprensen 7 physiol Chem 64 120 (1900) See also Northrop J Gen

Physiol 9 767 (1926)

Northrop (J Gen Physiol 9 767 (1926)) has improved the formal titration by titrating first to pll 7 with neutral red as the indicator then adding formaldehyde and titrating to pH 9 with phenolphthalein

colored to correspond with the color of the unknown solution. The accuracy of the formol titration is considerably improved by controlling the final pH adjustment and titration potentiometrically

Procedure. The solution to be analyzed, if carbonates, phosphates, and ammonla are absent, is made neutral to litmus (paper) and the solution titrated with formaldehyde as below st In case carbonates, phosphates, or ammonia are present a preliminary treatment is necessary which will vary according to the quantity of ammonia present

- (a) FOR SMALL AMOUNTS OF AMMOVIA Applicable to most urines Fifty mi of the material under examination are pipeted into a 100-ml measuring flask and I ml of phenolphthalein solutiones and 2 c. of solid barium chloride are added, the whole is shaken, to saturate the solution with barlum chloride, saturated barlum hydroxide solution is added until the red color of the phenoiphthaleln develops and then an excess of 5 ml 1s added The flask is filled to the graduation mark with water, shaken and permitted to stand for 15 minutes, after which it is filtered through a dry filter Place 80 ml. of the clear red filtrate (which corresponds to 40 ml of the liquid under examina tion) in a 100 ml. measuring flask, neutralize to litmus and dilute to 100 ml with freshly boiled water Equal portions of this solution, 40 ml (equivalent to 16 ml of the original solution), may be taken for analysis, one for the formol titration and the other for the determination of ammonia nitrogen
- (b) For large Amounts of Ammonia After the treatment with pheool phthalein, barium chloride, and barium hydroxide, and after the solution has heen diluted to 100 mi as in (a) above, the ammonia is distilled off, in vacuo

In case the solution is deeply colored, as in protein digests, it may be necessary to decolorize" before the tetration is attempted

Final Titration. For the final titration a volume of from 20 to 40 ml which contains approximately 25 mg of nitrogen is the most desirable A control solution is run composed of an equal volume of bolled distilled water and 20 ml of the formaldehyde mixture 12 This control solution is colored 13 50 that its tint matches that of the solution to be titrated.

To this control is added about half the volume of 0 2 N aikail which will be used in the titration of the solution under investigation and it is then titrated with 0 2 N acid to a faint red (first stage) 74

An additional drop of 0 2 N alkall is added, which imparts a distinct red to the solution (second stage)

- 47 As a standard of comparison the litmus paper used for neutralization is contrasted with a similar piece dipped in a phosphate solution having a neutral reaction (M/15 KH-PO₄ and M/15 \ailii O. in the proportion 40 60)
- 1 solution of 0.5 g of phenolphthalem in 50 ml of alcohol and 50 ml of water 19 The determination of ammonts may be dispensed with in case a separate determina-
- tion is made 16 For particulars with regard to the distillation etc. see Henriques and Sørensen
- 2 physiol Chem 64, 137 (1909) 11 For methods see Jessen Hansen Abderhalden a Arbeitsmethoden Vol 6 p 262 1912
- The formaldehyde solution is freshly prepared for each set of determinations as follows to 50 ml of commercial forn aldehyde (formol) (30 to 40 per cent) add 1 ml of the phenolphthalein solution 02 \ alkali is then added until the mixture acquires a faint red
- er lor The volume of the formaldehyde used will vary with the volume of the solution to he analyzed approximately 10 ml of the formalin se lution are added for each 20 ml of the unknown solution 14 Solution of Bismarck I rown is very satisfactory for urines. Tropeolin O tropeolin
- OO p-nitrophenol methal orange or alizarin sulfonate may be used
 - 1. It is procedure is recommended in or fer that the final volume of the control and it e unknown soluts as shall be approximately the same when the process is complete

The solution to be analyzed is now titrated to the color produced in the second stage of the control. The formaldehyde mixture is now added, 10 ml for each 20 ml of the solution, and the mixture again titrated to the second stage with 0.2 N nikali

Two drops of the 0 2 N alkali are now added to the control solution which assumes a deep red color (third stage) 9 2 N alkali is now added to the solution under examination until it assumes a color corresponding to the third stage of the control. This completes the titration

CALCULATION The calculations are similar to those which pertain to any acidimetry procedure Lach ml of an 0.2 \ alkalı or acid solution is convalent to 0.0028 g of nitrogen. An example will illustrate the procedure 40 ml of solution (16 ml of urine) required 5 10 ml of 0.2 N NaOH, control 0 10 ml of 0.2 N NaOH total required for amino acids 5 00 ml equivalent to 0 014 g of nitrogen Ammonia nitrogen in 16 ml of urine, 0 007 g of N Then 0 014 - 0 007 = 0 007 g amino acid nitrogen in 16 ml of urme

Interpretation. The excretion of total amino acid introgen by a normal adult, as determined by this procedure averages between 0 4 to 1 0 g per day or from 2 to 6 per cent of the total nitrogen. Free amino acid nitrogen (see previous procedure) is considerably less than this, ordinarily 0 5 to 1 0 per cent of the total mitrogen This discrepancy is due to the relative nonspecificity of the formal titration. For further aspects of interpretation, see preliminary discussion, and also previous procedure

3. Other Methods. Foliu proposed the colorimetric determination of amino acids in urine by reaction with naphthoguinone sulfonic acid.76 after preliminary treatment of the urine with Permutit to remove ammonia, which interferes Although this color reaction is sufficiently rehable for its use in the determination of blood amino acids where inter ference by ammonia does not arise, the procedure is not recommended for urine analysis since the Permutit treatment removes some of the amino acids as well as ammonia A more satisfactory procedure for removal of ammonia might render the method applicable to uring Albanese and Irby¹⁷ have proposed a simple titrimetric method for the determination of urmary ammo acid nitrogen, their procedure appears to have approvimately the specificity of the formol titration. For details of the nitrous acid reaction for urmary amino acids, see Van Slyke and Kirk 78

CREATININE

Folin's Method Principle This method is based upon the characteristic property possessed by creatinine, of yielding a certain definite color reaction in the presence of pierie acid in alkaline solution. This reaction (Jaffé reaction) is due to the formation of a red trutomer of creatinine picrate . The production of this tautomer is ' dependent upon the formation of a salt, a keto-enel change within the creatining molecule, and a change in the pieric acid molecule involving the hydrogens in the meta positions and, probably all three nitro groups 1 so

⁷⁴ This is best accomplished by adding alkali until the color is deeper than that of the control then acid again until lighter and finally alkali to the desired color

¹⁴ This procedure is described for amino scids in blood See p 565

^{**} This procedure is described for animo acids in blood See p. 505

**Albanes and Irby J Biol Chem. 152, 533 (1943)

**Van Slyke and Kirk J Biol Chem. 192, 601 (1933)

**Greenwald and Gross J Biol Chem. 59 601 (1924)

**Greenwald J Am. Chem Soc 47, 1443 (1923) J Biol Chem. 80 103 (1928)

In the original Folin macromethod, 4: 10 ml of urms are used for a determination, the color is developed at a final volume of 500 ml, and comparison is made visually against an artificial standard (0.5 N potassium bichramate solution) under rigorously defined conditions The Folin micro modification described here is preferred because it requires less urine, is based upon comparison against standard creatinine solutions, and is therefore more flexible and accurate, and is adaptable to either colorimetric or photometric measurement *2

Procedure.

- (a) FOR COLORIMETRIC COMPARISO. Measure 1 ml. of urine into a 100-ml volumetric flask, and in a second similar flask place 1 ml. of the standard creatinine solution,12 containing 1 mg. of creatinine per ml. To each flask add 20 ml. of 1 per cent plcric acid solution" (measured with sufficient accuracy from a graduated cylinder), followed by 1.5 ml. of 10 per cent sodium hydroxide solution. Mix gently and allow to stand 15 minutes. Dilute to the mark with water and mix by inversion. Compare standard against unknown In the usual way, setting the standard at 20 mm.
- (h) FOR PHOTOMETRIC MEASUREMENT Measure 0 5 ml. of urine into a 100-ml volumetric flask and add 0 5 ml. of water. In a second flask, which serves as a blank, place 1 ml of water. To each flask add 20 ml. of 1 per cent picric acid solution, followed by I 5 ml. of 19 per cent sodium hydroxide solution lik gently and allow to stand 15 minutes. Dilute to the mark with water and mix by inversion Transfer portions of blank and unknown to photometer cuvettes. Set the photometer to zero density with the blank, at 520 mu, and determine the density of the unknown. Alternately, the photometer may be set to zero density with water, and the densities of both blank and unknown determined The density of the blank is then subtracted from the measured density of the unknown, to obtain the true density of the unknown.

CALCULATION For colorimetric measurement

Reading of Standard | mg ereatinine = mg ereatinine in volume | mg treatinine in volume | furine analyzed

With a unne volume of 1 ml, and a 1-mg, standard the result of the calculation gives the creatinine content of the urine directly in mg per ml or g. per liter Urines containing 0 75 to 1 25 g. of creatinine per liter (i.e. the standard and unknown read within a few mm of cach other) may be accurately determined with the 1 mg. stand ard described For values outside this range the calculation is not accurate because of the deviation from Beer's law shown by the Jaffé reaction for creatinine In such cases the determination must be repeated using less or more urine as the case may be, to provide a portion of sample containing approximately I mg. of creatinine When this is done differences in volume between standard and unknown before adding the

[&]quot; Fulin J Biol Chem , 17, 469 (1914)

¹¹ for studies on the photometric determination of creatinine by the Jaffé reaction see Bousness and Taussky J Bud Chem 158 581 (1945) Peters J Biol Chem 146, 179 (1942) The use of 3,5-duntrobenzous and for the colormetric determination of creatmine has been proposed by Benedict and Behre J Biol Chem. 114, 515 (1936) and Langle) and Lvans shed 115, 333 (1936)

¹² Standard Creatinine Solution Dissolve 1 g. of pure dry creatinine in aufficient 0 1 by drochloric acid to make I liter and mix well. This solution contains I mg. of creatinine per ml. and is stable indefinitely Pure creatinine may be purchased from laborator) supply houses or may be made from urme by Benedict a method (see p 800) Creatinine zine chlorule may be used in place of creatinine in which case I 61 g, of the creatinine-rine chi stide salt is used per liter of 01 and to give the standard solution containing 1 mg of creatmins per ml

^{**} Prepared from purified partie acid (see Appendix). Keep in a dark bottle.

pieric acid must be equalized by the proper addition of water to one or the other as required. The further procedure and the calculations are as described. For very dilute urines (creatinine content less than 0.2 g per liter), the Shaffer modification of the Polin procedure may be used ⁴³

For photometric measurement the creatinine content of the unknown is established from its photometric density by reference to a calibration curve showing the densities for known amounts of creatinine determined by the procedure described. Direct calcu-

lation of results in terms of Beer's law is not valid in this procedure because of the lack of adherence to Beer's law over the concentration range in question

To prepare a calibration curve, measure 00, 025, 050 075, and 100 ml portions of standard creatinine solution. containing 1 mg per ml, into separate 100-ml volumetric flasks (preferably in duplicate) Add water, where necessary to bring each to 1 ml volume, and treat with pierie acid and alkali exactly as described under "Procedure for Photometric Measurement," above Determine the densities for each of the standards using the first or blank solution to set the photometer to zero density, or set the photometer to zero density with water and subtract the determined density of the hlank from that for each standard to obtain the true densities of the standards On a sbeet of ordinary cross-section paper, plot the density of each standard on the

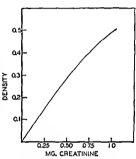


FIG 247 TYPICAL CALIBRATION CURVE FOR PHOTOMETRIC DETERMINATION OF CREATININE IN URINE

y axis against the amount of creatinine present, m mg, on the x axis. Draw a smooth curve to include all the points. A curve similar to that shown in Fig. 247 will be obtained

The amount of creatinine present in an unknown is obtained by reference to the curve If 0.5 ml of urine is used in an analysis, the result from the curve gives the creatinine content in this volume of sample. Multiply by 2 to obtain the creatinine content in fig. per ml or grains per liter.

The spectrophotometric characteristics of the creatinine color are shown in Fig. 248 At 520 ms and in a 1-cm cuseful the densities for various amounts of creatinine have approximately the values indicated in Fig. 247 Any creatinine content up to about 2 g per liter may be accurately determined under these conditions. For higher values or for photometric measurement at greater depth of solution the urms should be diluted, a portion of the diluted sample used for analysis and the creatinine content as read from the curve multiplied by the dilution to obtain the true creatinine content of the original undiluted urine

The sources of error in the use of a previously prepared calibration curve have been discussed in Chapter 23 For accurate results the curve should be chicked at intervals and reconstructed in necessary. In an analysis every effort should be made to have the conditions as nearly identical as possible with those prevailing at the time the curve was established, the time of standing (after adding the alkali and after

[&]quot;Shaffer J Biol Chem 18, 525 (1914)

diluting) and the room temperature are particularly important in this connection. The curve should be checked or reestablished with each new lot of pieric acid

Interpretation. The daily exerction of creatinine by normal adults ranges from about 10 to 18 g, under certain conditions it may be considerably higher. The value is nearly constant from day to day for a given normal individual, being influenced by the diet only to the extent that the diet itself contains significant amounts of creatinine, as in the case of a heavy meat diet. Creatinine exerction is not influenced by exercise or by the level of introgen metabolism in the body. Creatinine appears to be entirely a waste product, unutilizable by the body, and is exercted almost if not entirely quantitatively when ingested or injected. The relative constancy of creatinine excretion on a creatinine-free diet

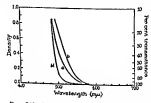


FIG 248 ABSORPTION SPECTRA OF COLORED SOLUTIONS OBTAINED IN URINARY CREATININE METHOD

For the alkalma pierate blank (bl) blank plus 0 2 mg. (a) and 0 50 mg. (b) of creatmine Solution depth 1 cm

appears to reflect some constant metabolic process involving the body creatine (from which creatinine is almost certainly derived), but the nature of this process is still obscure (see p 797 for further discussion) and therefore the significance of a constant creatinine excretion cannot be too precisely evaluated. Since the bulk of the creatine excretion cannot be too precisely evaluated. Since the bulk of the creatine of the body is in the muscles, there is a rough correlation between the creatinine excretion and the amount of muscular tissue in the body, for example, obese individuals excrete less creatinine relative to their body weight than do thin persons. The number of mg of creatinine excreted daily per kg of body weight is, hown as the creatinine coefficient in them of the number of mg of creatinine integer excreted daily per kg of body weight, the normal range in this case is 7 to 11.

Creatinine exerction is decreased in disorders associated with muscular atrophy and muscular weakness. It increases with increased tissue catabolism as in fever.

¹⁴ See Hobson Brochem J., 13 1425 (1939)

CREATINE

1. Folin Method.¹¹ Principle. Creatine on boiling with acid is transformed into creatinine. By determining the creatinine content before and after treatment with acid (pieric acid is used here), the amount of creatine present may be obtained by difference. The method is not applicable to diabetic urines, since acctooe and glucose interfere.

Procedure. 48 Measure the urine (I ml. for colorimetric measurement, 0.5 ml. for photometric measurement) into a 300-ml. flask, Add 20 ml. of 1 per cent picric acid solution and a few small pebbles or pieces of alundum. Weigh the flask and contents on a rough balance to the nearest gram. Add about 150 ml, of water and heat to boiling. Boil gently for 45 minutes, then more rapidly until the volume has been reduced to somewhat under the original volume of urine plus pieric acid solution, as established by weighing, Add sufficient water to the flash to restore the original weight, within about a gram Cool to room temperature (this is important). Add 1.5 ml, of 10 per cent sodium by droxide solution, and mix. Let stand 15 minutes. Rinse into a 100-ml. volumetric flask, dilute with water to the mark, and mix For colorimetric measurement, compare against a standard creatining solution, containing 1 mg. of creatinine, prepared in a second 100-ml, volumetric flask as described on p. 900 for the determination of urinary creatinine. For photometric measurement, determine the density (corrected for the picrate blank) as described on p. 900, and obtain results from a calibration curve, as described helow

CALCULATION For colorimetric measurement

Reading of Standard Heading of Unknown High Standard High

From this, the "total' creationse content per liter or in the 24-hour samplemay be obtained Subtract from the "total" creationse value the preformed creationse content, determined separately as described on p 900, to obtain the creatine content (expressed as creating)

(Total creatinine - Preformed creatinine) = Creatioe (as creatinine)

To convert creatine expressed as creatmine into the amount of creatine itself, multiply by 1 16

(Total creating - Preformed creating) × 1 16 = Creating

The same precautions concerning close agreement between standard and unknowo that are emphasized in the determination of urmary creatinine are applicable here. The total creatinine content of the sample should not be greater than about 1.25 g per liter, when a 1-mg standard is used. If it is higher than this, repeat the analysis using less urine, or an adiquot of diluted urine.

For photometric measurement from the determined density of the unknown (corrected for the pierate blank), obtain the 'total" creatinine content of the volume of urine used by reference to a calibration curve as described on p 901 for the photometric determination of urinary creatinine (the same curve may be used). Determine the creatine content as described above, by subtracting from the "total" creatinine

¹¹ lohn J Biol Chem 17, 469 (1914)

in The reagents required are those used for the determination of urmary creatingue (p. 900)

90 t

content the amount of preformed creatmine present determined separately. If the photo metric density of the vacaous is beyond the Limits of the calif ration curve, repeat the analysis in ng less time

Interpretation. Creatine occurs only in small amount (approxim. e range 0 to 200 mg. per day) in the urine of normal adults, but is found in larger amounts in the urine of children. More creatine is found in the urine during activity than when at reat. Hobson's found significant creatmuna (in one instance 14 g. per day) in the case of adult much athletes in training, on a high earbohydrate diet, decrease in earbohydrase intake resulted in a markedly decreased exerction of creatine. Albanese and Wangerin's found significant amounts of creatine in the urine of 30 normal adults, with no differences but veen sexes. According to these inve.ti, ators earlier viev a that creatinuria is not found in normal adults are possibly due to technical difficulties in the analysis

Increased creating exerction is noted in pregnancy in fasting, and after high water ingestion. A significant creatinum is noted in many pathological conditions associated with malnutrition and disintegration of muscular tissue etc Ingestion of creatine by normal adults does not increase the creatine content of the unne 10 the ingested creatine is completely retained. In certain pathological conditions however notably progres we muscular dy trophy this ability to retain in seted creatire is impaired at d extra creatine appears in the urine after a test dose This is the basis for the creatine tolerance test " which is sometimes used clinically for diagnostic purpose

URIC ACID

I Direct Colorimetric Method of Benedict and Franke ** Principle* The duted urme is treated directly with arsenophosphotungstic and reagent and sodium cynnide. The blue color obtained is compared with that of a standard uric acid solution treated in the same way. This method is known to be somewhat nonspecific for uric acid, as are all direct methods (see discussion under the other methods described in this section) but is believed to be quite satisfactory for many purposes.

Procedure 52 The urine 53 is so diluted that 10 ml will contain between 0 15 and 0 30 mg of uric acid (Usually a dilution of 1 to 20 is satisfactory) Team of the diluted urine are measured into a 50 ml volumetric flash, 5 ml of the 5 per cent sodium cyanide solution (poisonous') are added from a buret, followed by 1 ml of the arsenophosphotungstic acid reagent. The contents of the flash are mixed by gentle shaking, and at the end of 5 minutes diluted to the 50 ml mark with distilled water and mixed. For colorimetric measurement, this blue solution is then compared with a simultaneously prepared solution obtained by treating 10 ml of the standard uric acid solution (0.2 mg of uric acid) in a 50 ml flask with 5 ml of the sodium cyanide solution, 1 ml of the reagent, and diluting to the mark at the end of five minutes. For photometric measurement, determine the densities of unknown and standard at 520 m μ_1 setting the photometer to zero density with a blank solution obtained by treating 10 ml of water in a 50 ml flask with eyanide and reagent eractly as described for unknown and standard

CALCULATION For colorimetric measurement

Reading of Standard \times 0.2 \times $\frac{D}{10}$ = uric acid content of the urine in g per liter

D is the dilution (usually 20) of the urine For most neutrate results the standard and unknown should read within a few mm of each other. If they differ significantly repeat the analysis using a greater or lesser dilution of the urine as required.

^M Benedict and Franke J Biol Chem 52 387 (1922) For a modification of this method claimed to be superior see Christman and Raywitch J Biol Chem 95 115 (1932) \ photon etre version of the Christman Raywitch procedure is used in the uncase procedure of Buchanan Block and Christman described in the text (p. 907)

[&]quot;Solutions Required for first Verificial versions in the easy Control of the state

² Sodium Cyaride \ 5 per cent solution of sodium cyaride containing 2 ml of concentrated NH₄Off per liter which should be prepared fresh once in about 6 to " weeks

³ Uric Acid A standard solution of uric acid acidified with 13 drochloric acid containing 0.2 mg of uric acid in 10 ml is semployed. This solution may be readily prepared by dilution of Benedict's phosphato standard uric acid solution described in connection with the Newton uric acid in ethod for blood (seep 5.63). Vensure J0 if of the plo 1 hate standard solution (containing 10 mg of uric acid) into a 500-ml volumetric flask and dilute to about 400 ml with distilled water. 4d 2.5 ml of dilute 13 drochloric acid (made by diluting 14 olutions of 14 certon and diluted is solution to 500 ml and mix. This dilute attandard solution about d be prepared fresh from the pl osphato standard every 10 days to 2 weeks.

[&]quot;Une and tends to I recipitate out of urms on short standing I articularly if il e urms as concentrated or and 't all yes shoul it erefore be carried out as soon as I on ile after of taming it e urms and on it e well mused sample out crasse results will be maleading of taming it of the maleading of the male and on the well mused sample out crasse results will be maleading.

For photon etric measurement

Density of Standard \times 0.2 $\times \frac{D}{10}$ = uric acid content of the urine, in g per liter

As above D is the dilution of the urine. The spectrophotometric characteristics of the uric acid color are shown in Fig. 249. At \$20 mp and in a 1-cm cuvette, the stand and has a density of approximately 0.300. Agreement with Beer's law is satisfactory only at a unc acid content about equal to that of the standard For higher values or for photometric measurement at greater depths of solution carry out the analysis of a greater dilution of the urine. For more accurate results prepare a calibration curve as described for the following method (p. 908)

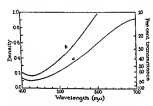


Fig 249 Absorption Spectra of Colored Solutions Obtained in Benedict Franke Method for Uric Acid in Units For standards containing 0.2 mg (a) and 0.4 mg (b) of uric acid Solution depth 1 cm

Interpretation. For adults on a mixed diet the average excretion of uno acid is about 0.7 g. It arises from the purines of ingested food (exogenous uric acid) and from purines derived from the body tissues by disintegration of nuclear material (endogenous uric acid). This distinction between two metabolic sources of urinary uric acid first postulated by Folin many years ago, appears to be confirmed by more recent work based on the use of isotopes. Fixogenous uric acid depending entirely upon the diet is greatly increased by the ingestion of purine-rich foods (meta, liver, sweetbreads etc.) and reduced to a very low level on purine-free foods e.g., milk eggs etc. (see table, Chapter 33). Endogenous uric acid is influenced by exercise and by the diet (protein foods particularly giving rise to increases). It appears to be partly the result of gastrointestinal secretory activity. On a purine free diet the average exerction is 0.1 to 0.5 g. On a high purine diet the uric acid output may be 2.g. per day

In gout the urne and content of the urne is low preceding an attack and increases during the attack this fall and rise being more or less characteristic The excretion rises after encopine administration, apparently because of increased kidney activity, and after the administration

^{**} Plentl and Schoenheimer J Biol Chem 153 203 (1944)

ACTH In leukema the excretion is extremely high due to nuclear destruction. The uric acid content of the urine is of importance in relation to the formation of uric acid calcul. The administration of alkali car bonates and citrates by decreasing the acidity of the urine increases its solvent power for uric acid, and decreases the possibility of the formation of this type of calculus.

2 Uricase Method of Buchanan Block, and Christman 100 Principle The color intensity resulting from the application of a une acid color reaction directly to a portion of the diluted urine is determined photometrically. According from the incubated with a preparation of the enzyme uncase which specifically destroys urine acid, and the uncase acid color reaction applied to this incubated sample. The difference in color value before and after treatment with uricase is considered to be a measure of the true urine acid content of the urine.

Procedure 101

(a) Total Colon Transfer 5 ml of urine to a 250 ml volumetric flask, dilute with water to the mark, and mix Transfer 10 ml of this diluted urine to a 50 ml volumetric flask and add 15 ml of water. In a second similar flask place 25 ml of water, this serves as a photometric blank. To each flask add 25 ml of urea cyanide solution, followed by 1 ml of arsenophosphotumistic.

100 Buchanan Block and Christman J Biol Chem 157 fS1 (1945) For a similar procedure see Schaffer J Biol Chem 153 163 (1944)

118 Reagents Required Urea Cyanide Solution Dissolve 25 g of pure sodium cyanide and 50 g of anity aforus sodium earbonate in 300 ml of water Cool add 70 g of pure urea and allute with water to 500 ml This solution is usable for several months even though a slight precipitate will settle out on standing. It is extremely posionous and must be handled carefully.

Arsenophosphotungsisc Acid. This is the same reagent that is used in the Benedict Franks method (see p. 905)

Urante Powder Remove the superficial (at from 5 pounds of fresh beef kidneys and grand in a meai-chopper Transfer to a large wide-mouthed bottle and wash by running in cold tap water slowly through a tube reaching to the bottom of the bottle until the supermistant fluid is quite clear and colorless Homogenize small portions of the resulting material in a Waring blendor with an approximately equal weight of benzere. To the combined total homogenize add 2 volumes of cold acctone. Allow the precipitate to settle filter through cheeseclot for a towel and squeeze dry. Suspend the solid material in about three times its weight of acctone allow it to settle and again filter off. Repeat until the resulting powder is thoroughly dehydrated and defatted Spread the material on towels and allow to dry in air overnight. Screen the powder through a 40-mesh sieve and store in a vacuum desirection.

This powder should give no blank color when carried through the incubation procedure and subsequent color reaction described in the text. To text for eartisty prepare a special lith ium carbonate per liter by following the procedure described on p 500 for the preparation of the Folin stock urns and solution up to but not including the addition of formalichy de and sulfure and which if present will interfere with urnses centrity. Each mil of this solution contains 1 mg of urns and incubate 3 to 4 ml of this solution with 250 mg of the urnsase powder borste buffer et and continue with the cofor reaction as for it e analysis of an unknown. If the urnsase is active no readual color will be obtained The special lithium carbonate standard is unstable and must be made up fresh on the day of use

Standar I Urie 1 end Sol tion. The Folin stock standard solution is used containing 1 mg of the acid per ml. prepared as described on p 550 in connection with the determination of time acid in bloof. Suitable dilutions of this stock standard are used in the preparation of the calif ration curve as described in the text. Dilute standards are unstable and n ust be prepared fresh out the day of use.

Borale Buffer (pil 9.2) Dissolve 12.4 g of borse acid in 1 liter of 0.1 \sodium hydroxide solution.

10 Per Cent Sods im Tungatate 0.66 \ Sulfure 4cid. The same reagents used in the preparation of Folia Wu blood filtrates (see p. 543)

acid. Both of these reagents are extremely polsonous; they should be handled with care, and always dispensed from burets. Immediately after the addition of the arsenophosphotungstic acid, dilute the contents of the flask to the 50-ml. mark with water and mix thoroughly. Allow to stand for exactly 30 minutes after so diluting (in serial analyses a suitable time schedule should be established), then determine the density of the unknown in a photometer at 690 mu, setting the photometer to zero density with the blank.

(b) RESIDUAL COLOR, Place 5 ml. of the original undiluted urine in a small flash, add 10 ml. of water and a few drops of 0.04 per cent thymol blue solution as indicator. Titrate with 0.1 N sodium hydroxide solution to a defioite blue tint. Record the amount of alkali required, and discard the mixture. Transfer a fresh 5-ml. portlon of the urine to a 50-ml. volumetric flask, and add the predetermined amount of alkali (without indicator). Add 250 mg. of urlease powder, 5 ml. of borate buffer (pll 9.2), and wash down the sides of the flask with 10 ml. of water. Place in a water bath at 45° C. for 2 hours. Add 1 ml. of 10 per cent sodium tungstate and 1.5 ml. of two-thirds normal sulfuric acid. Dilute to 50 ml. with water, mix well, and pour ooto 2 dry filter. Transfer 10 ml. of the filtrate to a 50-ml. volumetric flask, add 15 ml. of water, and continue with the addition of the color reagents and measurement exactly as described above for "Total Color" procedure. The same reagent blank may be used for setting the photometer to zero density, or 2 5-ml. portion of water may be treated with uricase, buffer, etc., as described for the analysis of the urine, and the final solution after treatment with the color reagents used as a photometric blank.

CALCULATION

a Total Color Value Determine the amount of uric acid in mg equivalent to the photometer reading of the total color sample by reference to a calibration curve (see below) Multiply this by 5 (since the 10 ml of urine diluted 1 50 represents 0 2 ml of original urine) to obtain the uric acid content for the undiluted urine equivalent to the total color, in mg per ml (or g per liter)

b. Residual Color Value In a similar way determine the uric acid content equivalent to the resulual color, the value in mg obtained from the calibration curve gives directly the residual color equivalent of the original urine in mg per ml, or g per

liter, since 10 ml of urine dduted 1 10 are used

c True Uric Acid Content Subtract the 'residual color" value from the "total color" value, to obtain the true unic acid content of the urine

> Total Color - Residual Color - True uric acid content (as uric seid) (as urie acid) (in g per liter)

Multiply the result by the urine volume (expressed in liters) to obtain the uric acid ontent of the entire sample

Other dilutions and aliquots than those specified may be used if the final colored solutions are either too light or too dark for accurate photometric measurement, in such case the calculations must be corrected accordingly

The use of a calibration curve prepared from standard uric acid solutions is recommended because of the deviation from Beer's law shown by lower concentrations of uric acid To prepare such a curve, proceed as follows Dilute I mi of the stock Folin une acid standard (containing 1 mg per ml) to t00 ml with water and mix, thus obtaining a solution containing 0 0t mg of uric acid per mi Measure aliquots of this dilute standard into 50-ml. volumetric flasks (preferably in duplicate) to give a series of solutions of known uric acid content covering the range of satisfactory photometric measurement (0 0t to 0 12 mg for measurement at 2 cm solution depth or its approximate equivalent, as with the Livelyn photometer, 0 02 to 0 24 mg if 1-cm. cuvettes or their equivalent are to be used) Include a blank flask containing water alone in the series Adjust the volume in each flask to 25 ml by adding water where necessary, then add 25 ml of urca-cyanide solution to each flask. At timed (e.g., one-minute) intervals add I inl of arsenophosphotuogstic acid to a flask dilute immediately to 50 ml with water and mix thoroughly At the end of 30 minutes, set the photometer to zero density with the blank and determine the density for each standard exactly 30 minutes after diluting and mixing Plot the determined densities against the amount of uric acid present (in mg) on cross section paper, and draw a smooth curve to include the points

In an analysis, the determined density of an unknown is translated into its equivalent uric acid value by reference to the curve. For accurate results the curve should be checked at intervals particularly if new reagents are made up, and reconstructed if necessary In an analysis environmental and other conditions should reproduce as consistently as possible those prevailing at the time the curve was established. For further discussion concerning the validity and use of calibration curves in photometric analysis see the section "Photometry in Chapter 23 p 512

Interpretation. Results by the uncase method described here indicate that only 80 to 90 per cent of the color obtained by direct treatment of urine with uric acid color reageuts may actually be due to urie acid, this discrepancy may be much greater if the diet contains significant amounts of the methyl vanthines (eaffeine, theophylline, and theobromine, found in coffee, tea, and cocoa) In such justances, the uriease method appears to be even more specific for une acid than the "isolation" procedures (see following method) For further aspects of interpretation, see under previous method

3 Colorimetric Methods of Folin 102 Principle Phosphotungstic acid is reduced by uric acid with the production of a blue color which is compared with that produced with a standard solution of uric acid Polyphenols also react, thus giving too high results, and amino acids decrease the color The direct application of the reaction to urine is oot therefore an accurate method but may he useful for many clinical pur poses The indirect method in which the uric acid is first separated by precipitation with silver nitrate is claimed to be an accurate method if conditions are strictly followed

Procedure

(a) INDIRECT METHOO Half fill a 160 ml volumetric flask with water With a Folin Ostwald pipet introduce 1 ml of the urine Add 10 ml of the chloride acetate solution 102 and then without shaking so as to avoid foaming, dilute to the mark with water and mix

Transfer 5 ml of the diluted urine to one 15 ml centrifuge tube and 3 ml plus 2 ml of water to another To each add 3 ml of the sliver nitrate solution

Folm J Biol Chem 101, 111 (1933) 106 311 (1934)
 Chloride Accide Solution, A solution containing 1 per cent NaCl 2 per cent cryst.

sodium acetate and I volume per cent of 99 per cent acetic acid 5 Per Cent Solution of Silver Astrote This solution even if perfectly clear when first prepared may develop a slight color on standing Tl is color is most quickly produced by heating to 100° C for 2 hours in a flash covered with a beaker. After cooling add a few ml of a solution containing 50 mg of NaCl shake thoroughly and filter through a double layer of quantitative filter paper until crystal clear Thereafter the solution will rerperfectly colories and need not be kept in brown bottles.

Otler solutions are the same as those used in blood methods (see p. 560)

and centrifuge at once fairly rapidly for 4 to 5 minutes so as to get perfectly clear supernatant solutions. A few tiny flakes may float on the surface but these contain no uric acid. Decant and drain over a sink. It is permissible to let cold tap water rinse the mouths of the tubes during the draining. With a 25-ml cylinder or a buret, to each tube add i0 ml of the urea cyanide solution (poironous!) Stir immediately and simultaneously with glass rods until the two sediments have completely dissolved. Transfer the silver cyan ide solutions to test tubes graduated at 25 ml and rinse with exactly 5 ml of water. Nix by whirling at an angle of about 60° until the solutions are visibly uniform. In another graduated test tube place 5 ml of the standard uric acid solution containing 0.02 mg of uric acid together with 10 ml of the urea cyanide solution and mix

With a 10 ml blood pipet add to each of the three tubes 4 ml of the unit and reagent and let atand for 20 minutes Dilute to volume and mix For colorimetric measurement, make the comparison between the standard and the unknown which is nearest to it in depth of color. When the staodard is set at 20 mm, colorimetric readings between 35 mm and 10 mm are at ceptable. For photometric measurement, only one unknown is necessary Determine its density, and that of the standard, in a photometer at 60 mm, setting the photometer to zero density with a blank prepared by treating 5 ml of water with urea cyanide and color reagent, etc., exactly as described for the preparation of the standard.

(b) Direct Method Half fill a 100 ml volumetric flask with water With a Folin Ostwald pipet introduce 1 ml of urine, dilute to volume and mir Introduce into test tubes graduated at 25 ml, 5 ml of the diluted urine and 3 ml of the diluted urine med 3 ml of the diluted urine plus 2 ml of water To another graduated test tube add 5 ml of the standard uric acid solution Add 10 ml of the urica cyanide solution (Doutonour') to each, mix, and add 4 ml of the uric acid reagent Let stand for 20 minutes Dilute to volume and mlx Continue with colormetric or photometric measurement as described above for the indirect method

CALCULATION For colorimetric measurement

 $\frac{\text{Reading of Standard}}{\text{Reading of Unknown}} \times 0.02 \times \frac{100}{\sigma} = \frac{\text{Uric aci 1 content of indiluted}}{\text{urine in g per liter}}$

 σ is the volume of $d\bar{u}uted$ urane (o mi) or 3 ml) used in an analysis. The same calculations are used for both indirect and direct methods

For photometric measurement

Density of Luknown Density of Standard $\times 0.02 \times \frac{100}{e}$ = Une acid content of undiluted urine in g per liter

The significance of σ is the same as for colorimetric measurement. The conditions for satisfactory photometric measurement are the same as those given on p -ofi in connection with the determination of une acid in blood, since the same standards and color reaction are used.

Interpretation. See previous methods

4 Folin Shaffer Method Principle Ptosphates and some organic matter are removed by means of uranium acetate. The unc acid is precipitated as ammonium urate which is titrated with potassium primanganate.

¹³⁴ Folm and Shaffer Z physiol Chem 32, 552 (t-201) This method was described in the 11th and previous editions of this book.

PURINE BASES (INCLUDING URIC ACID)

1. Kruger and Schmid's Method: Principle. This method serves for the determination of both une acid and the purine bases. The principle involved is the precipitation of both the une acid and the purine bases in combination with copper oxide and the subsequent decomposition of this precipitate by means of sodium sulfide. The une acid is then precipitated by means of hydrochloric acid and the purine bases are separated from the filtrate in the form of their copper or silver compounds. The nitrogen content of the precipitates of une seid and purine bases is then determined by means of the Kieldahl method (see p. 874) and the corresponding values for une acid and purine bases calculated. For a study of this method as applied to the determination of purines in protein free tissue extracts, on a microscale, see Hitchings and Fiske. 188

Procedure, To 400 ml, of alhumin-free urine104 in a liter flask, 107 add 24 g of sodium acetate, and 40 ml of a solution of sodium hisulfite, 188 and heat the mixture to bolling Add 40 to 80 ml 303 of a 10 per cent solution of copper sulfate and maintain the temperature of the mixture at the holling point for at least 3 minutes. Filter off the flocculent precipitate, wash it with hot water until the wash water is colorless, and return the washed precipitate to the flask by puncturing the tip of the filter paper and washing the precipitate through by means of hot water Add water until the volume in the flash is approximately 200 ml, heat the mixture to boiling, and decompose the precipitate of copper oxide by the addition of 38 ml of sodium sulfide solution 116 After decomposition is complete, the mixture should be addified with acetic acid and heated to boiling until the separating sulfur collects in a mass. Filter the hot fluid with the ald of a filter pump, wash with hot water, add 10 ml of 10 per cent hydrochloric acid, and evaporate the flitrate in a porcelain dish until the total volume has been reduced to about 10 ml. Permit this residuo to stand about 2 hours to allow for the separation of the uric acid, leaving the purine bases in solution. Filter off the precipitate of uric acid, using a small filter paper, and wash the uric acid, with water made acid with sulfuric acid, until the total volume of the original filtrate and the wash water aggregates 75 ml Determine the nitrogen content of the precipitate by means of the Kjeldahl method (see p. 874), and calculate the uric acid equivalent.111

Render the filtrate from the uric acid crystals alkaline with sodium hydroxide, add accelc acid until faintly acid, and heat to 70° C. Now add 1 ml. of a 10 per cent solution of acetic acid and 10 ml. of a suspension of manganese dioxide¹¹¹ to oxidize the traces of urle acid which remain in the solution

¹⁰³ Hitchings and Tiske J Biol Chem , 149, 491 (1941)

¹⁰⁴ If albumus is present, the urine should be heated to boiling acidified with acetic acid and filtered

¹⁹⁷ The total volume of urino for the 24 hours should be sufficiently diluted with water to make the total volume of the solution 1600 to 2000 ml

¹⁰⁴ A solution containing 50 g sodium bisulfite in 100 ml water

The exact amount depending upon the content of the purme bases

He This is made by saturating a I per cent solution of sodium hydroxide with hydroken sulfation as and adding an equal volume of I per cent sodium hydroxide Ordinarily the addition of 30 ml of this solution is sufficient but the presence of an excess of sulfide should be proced by adding a drop of lead acetate to a drop of the solution Under these conditions a dark brown color will show the presence of an excess of sodium sulfation.

^{11.} This may be done by multiplying the nitrogen value by three and adding 3.5 mg to the product as a correction for the nne acid remaining in solution in the 75 ml

he Made by heating a 0.5 per cent solution of potassium permanguage with a little alcohol until it is decolorized

Agitate the mixture for 1 minute, add 10 mi. of the sodium bisuifite solution. and 5 mi. of a 10 per cent solution of copper suifate, and heat the mixture to boiling for 3 minutes. Filter nil the precipitate, wash it with bot water, and determine its nitrogen content by means of the Kjeldahi method (see p. 874). Inasmuch as the composition and proportion of the purine baser present in urine is variable, no factor can he applied. The result as regards these bases must therefore be expressed in terms of nitrogen.

Benedict and Saiki report cases in which the total purine nitrogen by this method was less than the uric acid nitrogen as determined by the Folia-Shaffer method. The inaccuracy was found to lie in the Kruger and Schmid method. To obviate this they advise the addition of 20 mi of glacial acetic acid for each 300 mi, of urine employed, the acid being added before the first precipitation.

Interpretation. The amount of purme hases excreted by a normal man is small and variable Values from 16 to 60 mg have been found. The purme base introgen is of course only a fraction of this. The amount excreted is influenced by the duet somewhat in the same way as is the excretion of une acid, heing also increased in disorders associated with increased une acid excretion such as leukemia. The purme bases form a higher percentage of the total purme excretion in the case of the monkey, sheep, and goat than in the case of man

2. Hunter and Givens' Modification of the Kruger-Schmid Method in Principle The Kruger-Schmid process is combined with the colonmetric method for unpaged (see p. 909)

Procedure. The first copper purine precipitate as obtained in the KrügerSchmid procedure is suspended in ahout 200 mi of water, to which there
is added about 1 mi of concentrated bydrochloric acid The muture is
vigorously holled, whereupon the whole or greater part of the precipitate goes
into solution Removal of the copper is effected by treatment, while hot, with
hydrogen suifide and excess of the suifide is completely expelled by renewed
boiling. Filtration under suction, and thorough washing of flask and filter
result in a fittrate which is perfectly clear and nearly colorless. This is concentrated if necessary, and made up in a convenient volume which must of
course be sufficiently large to retain, when cool, the uric acid in solution
Of this an aliquot part is utilized directly for the colorimetric determination
of uric acid. In the remainder the residual uric acid is destroyed and hase
determined according to the regular Krüger-Schmid procedure. This modification is recommended particularly where the amount of uric acid present
is minute.

3 Welker's Modification of the Methods of Arnstein and of Salkowski Principle. The phosphates are removed by treatment with magnesia mixture. The purine bases and uric acid are then thrown down as their silver salts and the nitrogen content of this precipitate determined.

¹¹² To dissolve the excess of manganese dioxide

¹¹⁴ Hunter and Givens J Bud Chem 17 37 (1914) 115 Dittman and Welker V Y Med J 89, 1134 (1909)

ALLANTOIN

1. Method of Larson 111 Principle The utme is treated with phosphotungstic acid to remove interfering substances Basic lead acctate is then added to remove excess phosphotungstic acid and residual interfering substances Excess lead is removed by sulfuric acid and excess acid neutralized by sodium hydroxide. The solution is then boiled with Folin ammoniacal reagent and acid molybdate reagent added to the cooled solution. After proper dilution a colorimetric comparison is made against a ling allantom standard.

Procedure Transfer 1 5g of phospbotungstic aciding to a 50-ml centrifuge tube and add 5 ml of water Rotate gently to insure solution, then add 5 ml of animal urine Centrifuge Immediately, place the tube in a refrigerator for one-half hour, then centrifuge again until perfectly clear. The addition of a crystal of phosphotungste acid should not cause further precipitation. Add 5 ml of basic lead acetate solution which precipitates the excess phosphotungstic nich as well as the remaining interfering substances. Centifuge the mixture, then treat with 5 ml of 5 per cent sulfuric acid to remove the excess lead, and centrifuge until perfectly clear. Pipet 2 ml of the resulting water clear liquid into a Folin-Wu sugar tube, neutralize with 5 per cent sollium hydroxide, and then indd 2 ml of Folin immoniacal copper reagent. The Place the tubes in a rapidly boiling water bath for 10 minutes, cool, then add 2 ml of acid molybdate reagent. The Dilute the tubes to volume and read in n

INLLargon J Biol Chem 94 727 (1932) According to Young and Conway (J Biol Chem. 142, 839 (1942)) the Larson procedure although giving reasonably good recovery of added allantom gives somewhat high results on urine compared to the procedure thoy describe which is claimed to be more specific and satisfactor. For details see the original artiele For a photometric version of the Young Conway procedure and method for the determination of allanton in blood see Young MacPherson Weatworth and Hawkins J Biol Chem. 152 245 (1944)

iii) Phosphos/4-tungstic Acid Dissolve 100 g of Na WO 2H O in about 100 ml of water with the aid of heat 4 did 10 ml of 85 per cent HaPO, and then 80 ml of concentrated HCI Cool After 4 hours or more fifter on a Buchner funnel and suck as dry as possible. Redussolve the precipitate in 120 ml of HoP pour the solution into a hier separatory funnel add about 90 ml of ether and then add 40 ml of concentrated HCI Shake After standing a few minutes there should be three layers of heund The lowest layer contains nearly all the complex acid If there are only two layers more ether must be added and the mixture of the complex acid If there are only two layers more ether must be added and the mixture of water and shake vigorously then add 50 ml of ether and finally 50 ml of concentrated HCI After standing the lowest layer it is should be perfectly elear is transferred to a crystallizing dish 4dd 30 ml of H₂O and 1 drop of liquid bromine and evaporate on a steam bath. The solution should be greenash in color II the slightest trace of dust or organic matter is present a pinksib color develops and 1 or 2 drops more of liquid bromine must be added to oxidize this foreign maternal.

Evaporate on the steam bath until crystals began to form on the surface. Let stand overnight. The crystals obtained are sucked as dry as possible on a large Buchner funnel liter sur-drying for one week, powder the crystals and keep in an amber glass container. This phosphotungstic acid should dissolve instantly to give a perfectly clear practically coloriess solution.

[&]quot;Felin Ammonuccal Copper Selation Dassolve 100 g of ammonuum sulfate in about 400 ml of water and filter into a volumenten letter flaak. 100 ml of 10 per cent sodium hydroxide are then added 12 g of sodium tartrate and finally a solution of 5 g of copper sulfate Dilute to volume and mix This research will not give a blank for months if kept in the dark in well filled tightly stoppered amber glass bottles. The bottles should be of small volume.

¹ Folin Acid Molybdate Reagest Prepare a stock solution of 30 per cent brominated sodium n olybdate as follows Dissolve 300 g of sodium molybdate in water and I make up to 1 liter The solution is slightly turbid 14d2 or 3 drops of hquid bronne and let stand

colorimeter against a 1 mg allantoin standard 120 Photometric data on this procedure are not available

CALCULATION

 $\frac{\text{Reading of Standard}}{\text{Reading of Linknown}} \times 1 \times \frac{100}{9} = \text{mg allantom per 100 ml}$

Interpretation. Allantom is found in only small amounts in human urine (35 to 45 mg per day), and appears to be mainly, though not entirely, evogenous in origin. It forms however, the principal end product of the purine metabolism of practically all mammals other than man and the anthropoid apes with the notable exception of the pure bred Dalma intain coach hound which exercise a considerable fraction of its purine mitrogen as uricacid. Thus over 90 per cent of the purine-allantom nitrogen exerction of the dog the cow, and the pig occurs as allantom. In these animals its origin is from exogenous and endogenous purines, and its exerction is influenced by much the same factors as is that of uricacid in man. It appears to be entirely a waste product since if nigeted into the blood of man or dogs it is exercted almost quantitatively in the urine.

HIPPURIC ACID

I Method of Griffith 13 Principle Ilippuric acid is extracted from urine subther in a continuous extraction apparatus. The residue obtained by distilling off the ether is treated with I romine and sodium hypobromite to destroy traces of urislippuric acid introgen is then determined in the residue by it is kijeldahl procedure.

supports acid introgen is then determined in the residue by it is Ajedian processing. In Quick is netted off-the hippure acid is extracted in a similar manner and the amino introgen determined by the formol turation (see p. 8.77). The older methods in whole displayds sof the hippure acid and turtation of the liberated beazons acid. These may give muleading results however because benzon acid may be present also in the form of electromic acid monoleance the charge is fluencing as supported in the processing the

CALCULATION If y represents the number of ml. of 0.1 N acid neutralized by ammonia in the Kieldahl titration, then

$$y \times 1.4 \times \frac{179}{14} = \text{mg. hippuric acid in 10 ml. urine}$$

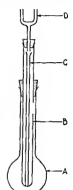
Interpretation. The average excretion of hippuric acid by a normal adult man is about 0 7 g. per day. The amount is increased by the inges-

tion of benzoic acid or fruits such as plums, prunes, and cranberries which contain, in addition to benzoic acid, certain other precursors of hippuric acid (quinic acid, etc.). It arises in part apparently from putrefaction products formed in the intestine. In herbivora it is often the most abundant nitrogenous constituent of the urme

2. Hippuric Acid Test for Liver Function (Quick125); Principle, If benzoic acid is ingested or injected, a major portion of it combines with glycine to form hippuric acid which is then excreted in the urine (for reactions, see p. 440) In man (but not necessarily in other animals) this synthesis appears to take place primarily in the liver if the liver is damaged, the amount of hippuric acid excreted is diminished relative to that found normally. It is claimed that chinically the decrease in hippuric acid excretion under test conditions is fairly proportional to the extent of hepatic impairment

Procedure. Just before the test, which is preferably given in the morning about I hour after a light breakfast (cereals or toast, and coffee, tea, or milk), the patient is instructed to empty his bladder as completely as possible, and the urine obtained is discarded. A dose of 6 g. of sodium benzoate dissolved in 30 ml. of water (flavored with oil of peppermint if desired) is then ingested,124 followed by half a glass of water. More water may be taken during the test if necessary but excessive water intake should be discouraged in order to keep the urine volume down. A complete specimen of urine is collected each hour after the test dose, for 4 hours.

Measure the volume of each hourly specimen in a graduate. If a specimen volume exceeds 150 ml., it is advisable to acidify slightly with acetic acid and to concentrate the sample on a water bath to about 50 ml. Combine the four specimens in a graduated cylinder, measure the volume, and transfer to a beaker. Add solld ammonium sulfate in the proportion of 5 g. for each 10 ml. of



250 CONTIN-UOUS EXTRACTION APPARATUS (GRIF-FITH).

.l. 500-ml. Kicldahl flask; B, glass tube (420 × 15 mm.) with side opening; C, glass tube (400 X 6 mm) with widened top and small openmgs at bottom; D, condenser.

urine, and stir to dissolve. Filter or centrifuge. To the clear filtrate or centrifugate in a beaker add sufficient concentrated hydrochloric acld (usually about I ml. is required) to render distinctly acid to Congo red or thymol blue (pll about 2).127 Stir vigorously with a class rod, scratching the sides of

in Indicator paper may also be used.

in Quick: Am J. Clin. Path , 10, 222 (1940). See also Weichselbaum and Probitein: J. Lab Clin Med. 24, 636 (1935-1939). Hepler and Gurley abid., 27, 1593 (1941-1942). 110 For details of the intravenous test, see Quick (loc. cit.).

the beaker to promote crystallization, then place in the refrigerator or in ice water for 30 minutes Fliter off the crystalline hippuric acid by suction on a small Buchner funnel, and wash the precipitate with several small portion of ice cold distilled water, using the wash water to complete the transfer of the crystals from the beaker to the Buchner funnel

Quantitatively transfer the precipitate and filter paper to a beaker (a little water may be used to aid in the transfer), add as unficient water to core, and heat to dissolve Add a few drops of phenolphthalein solution and titrate with 0 5 N sodium hydroxide solution to a permanent pink Record the burst reading

CALCULATION Lach ml of 0 a N sodium hydroxide is equivalent to 0 0895 g hippure and Therefore

ml 05 \ alkalı used \ 00895 = g hippuric acid titrated

To determine the amount of hippuric acid exercted a correction for the solubility of the compound must be added. In the presence of ammonium sulfate as described the solul dity has I cen estal lished as 0 001 g per inl. of urine. If λ is the total volume of urine in milliliters before a king the ammonium sulfate the solubility correction is $V \times 0$ 001 g. Therefore

(ml of 0 5 N alkalı used × 0 0895) + (V × 0 001) = g hippuric acid excreted
G hippuric acid × 0 68 = g benzoic acid

Interpretation Under the conditions of this test the average healthy adult will exercte 30 to 35 g (or even more) of benzoic acid in the form of hippunc acid. There is some indication that increased exerction above 3 g is roughly proportional to body weight (or surface area) in normal individuals ¹¹⁸ Any excretion of 27 g or more is considered nonpathological. Marked diminution in output is found in various liver disorders. For further aspects of clinical interpretation see texts on clinical diagnosis.

LACTIC ACID

Method of Friedemann and Graeser "Principle By treatment with phosphoric acid and potassium permanganate the lactic acid is converted to acctaldeby de

CII, CHOH COOH → CII, CHO + CO + H,O

The aldehyde is bound with sodium bisulfite

CH₁CHO + \allSO₂ → CH₁CHOH SO₂\a

The bound suif te is titrated indirectrically

Procedure A 10 to 25 ml sample of urine is introduced into a 250 ml volumetrle flask, 10 ml of 26 per cent CuSO, 511,0 and 10 ml of a suspension of Ca(011); are added "The suspension is diluted to the mark and filtered, an allquot representing not more than 5 ml of urine beling used for analysis

tations of the method for lood (see p. 624) to lk culture med a sid thouse supported leagents. Calcum Hydroxide Suspension 1 kg. of fresh unalsked time is alsked with water and immediately afterward sufficient water in added to bring the volume.

¹⁰⁸ Hepler and Gurley loc cut
¹⁰⁸ Friede nann and Graeser J Biol Chem. 100, 291 (1933). The authors describe adaption.

Ten mi of HiPO:-\inSO; reagent and a pinch of finely powdered taic are placed in a 300-ml Kieldahl flask, followed by the sample of urine About 85 ml of water are added and the flask attached to the apparatus (see Fig. 251) Ten ml of NaHSO2 solution are placed in the 150-ml extraction flask (receiver). The microburner is adjusted to bring the solution to boiling in about 3 minutes. Addition of oxidizing agent is begun as soon as vapors anpear in the condenser. The rate of addition is unimportant, but it is essential to have an excess of oxidizing agent throughout the distillation, as indicated by a brownish (not gray) color. It is best to add from 25 to 40 ml over a 15minute oxidation period. Shortly before the end of the oxidation the receiving flask is detached from the stopper and lowered. The glass tip is rinsed and the flask containing a total volume of 50 to 75 ml is cooled prior to titration

For the removal of excess bisulfite 1 ml of starch solution is added, followed by a sight excess of the strong lodine solution which is immediately removed by the cautious addition of 0 1 N thiosulfate. The walls of the flask are now washed down by a thin stream of water after which the end point is adjusted to a faint blue with dilute jodine solution

The flask is cooled and approximately 15 ml of saturated NaHCO, are added The solution is titrated with dilute standard jodine solution which is run in rapidly so as to keep pace with the decomposition into aidehy de and

to approximately 5 liters. The suspension is thoroughly shaken allowed to stand for a few seconds and decanted from the coarser particles.

Theorems and decinted from the Goarser particles and the Goarser particles to about 1000 ml and the solution is again well mixed. The reduction is complete in 5 to 10 minutes

Sodium Bisulfite 25 g are dissolved in 2 liters of water. The solution should be kept in a stoppered bottle

Starch Indicator 5 g of arrowroot starch are suspended in 10 to 20 ml of cold water and poured into 500 ml of boiling water, 500 ml of hot water are added and boiling is con tinued for 15 minutes. The flash is covered with a beaker cooled and kept in the refrigerator The supernatant clear solution only is used The solution will keep several weeks if care is taken to avoid bacterial growth

Strong Iodine Solution 40 g of iodine and 75 g of KI are dissolved in a small quantity of water and the volume is brought up to about 2 liters

Standard Iodine Solution The weak todine solutions (0 01 to 0 002 N) may be prepared either by dilution of a standard todine solution or by liberation of todine from 0.1 N todate the control of a standard forms solution or of the permanence of iodate solutions of the permanence of iodate solutions of the permanence of iodate solutions of the the permanence of iodate solutions of the third that the permanence of iodate solutions of the third that the permanence of iodate solutions of the third that the permanence of iodate solutions in the third that the permanence of iodate solutions is the third that the permanence of iodate solutions is obtained to the iodate of the third that the third that the permanence of iodate of the permanence of iodate of the permanence of iodate of the permanence of iodate of the permanence of iodate of the permanence of iodate of the permanence of iodate of the permanence of iodate of the permanence of iodate of the permanence of iodate of iodate of the permanence of iodate iodate of the permanence of iodate iodate iodate iodate of the permanence of iodate iod

to the mark Since iodine is volatile even from dilute solutions they should be kept cool Lactic Acid Standard. Lithium lactate is preferred because this salt is anhy drous and not hygroscopic It is prepared as follows

USP lactic acid (85 per cent) is diluted with an equal volume of water and a few drops of phenol red indicator are added Saturated (approximatel) 20 per cent) lithium by droxide or LiCO: (the former is preferred) colution is added to slight excess as is indi cated by the phenol red The solution is heated to boiling and the alkali is again added to slight alkalimity It is now cooled Four volumes of 95 per cent alcohol are added and after cooling for some time the mass of crystals is filtered off on a Buchner funnel and washed thoroughly with 95 per cent alcohol. This preparation is recrystallized from water an I d red at 100° C

⁹ to g of hithum factate are transferred to a 1 hier volumetric flask 1 nough H_2 :04 is added to bring the final concentration to 0.2 % when diluted to the mark. This 0.1 M lactic acid standard will keep for at least one year provided it is kej t away from strong light or stored in a refrigerator

Chap 31

hisulfite When this slows up 1 ml of 10 per cent Na₂CO₄ is added until the

end point persists for at least 12 minute Blank determinations should be run with filtrates prepared with the precipitating reagent and the CuSOs and lime reagents

CLICLIATION Lach ml of 0.01 N sodine solution used in the titration of bound sulfite is equivalent to 0.45 mg of lactic acid

Interpretation The lactic acid concentration is variable depending upon the volume from 4 mg in very dilute urine to as high as 25 mg per 100 ml in concentrated urine These values are increased after severe-exercise and probably in such pathological conditions as are accompanied

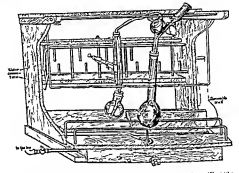


Fig. 251 Apparatus for the Determination of Lactic Acid (Friedle Many and Graeser)

by defining a substitute metabolism. For a discussion of lactic acid in the name see p. 812.

7

and Browne¹³² have described a quantitative photometric version of the naphthoresoccinol test (see p 843) which appears to give good results

GLUCOSE¹²⁴

1. Benedict's Method 115 Principle Benedict's reagent for the estimation of reducing sugars contains potassium thooyanate as well as copper sulfate, and in the rescace of the former a white precipitate of cuprous thooyanate is formed in reduction instead of the usual red precipitate of cuprous oxide. The small amount of potas sum ferrocyande also aids in keeping cuprous oxide in solution. As the precipitate formed is white the loss of all blue that in the solution, indicating complete reduction of the copper, is readily observed. The alkali used is sodium carbonate, which has the advantage over hydroxides of being less likely to cause destruction of small amounts of sugar. The solution also has the great advantage of being stable for an indefinite length of time. The method is recommended for its supplicitly and accuracy.

Procedure The urine, 10 mi of which should be diluted with water to 100 mi (unless the sugar content is believed to be low, when it may be used undiluted), is placed in a 50 ml buret and the volume adjusted to the zero mark Twenty five ml of the reagent 136 13 measured with a pipet into a porceiam evaporation dish or casserole (100 to 125 mm in diameter). 20 c of crystallized sodium carbonate (or one half the weight of the anhydrous salt) is added, together with a small quantity of powdered pumice stone or tal cum, and the mixture heated to boiling over a free flame and stirred with a glass rod to aid in dissolving the bulk of the carbonate. The diluted urine is now run in from the buret rather rapidly, until a chaik-white precipitate forms in noticeable amount and the blue color of the mixture begins to lessen perceptibly, after which the solution from the buret must be run in a few drops at a time, until the disappearance of the last trace of blue color, which marks the end point The final color at the end point may be yellow or brown, owing to urmary pigments, but there should be no trace of blue (or green) color The solution must be kept vigorously boiling and be stirred continuously throughout the entire titration If the mixture becomes too concentrated during the process, water may be added from time to time to replace the volume lost by evaporation. Any material which dries out on the sldes of the dish during the titration must be pushed back into the solution with the stirring rod before the end point is reached

¹¹¹ Maughan Evelyn and Browne J Biol Chem 126 567 (1938) See also Deichmann J Lab Chin Wed 28 770 (1943)

¹¹⁴ The method for sugar adopted by the Committee on Urinary Impairments of the Association of Life Insurance Medical Directors of America is given on p 923

With 10 and of heat dissolve the carbonate extrate and thoo; nanto in enough water to make about 500 ml of the maxture and filter if necessary. Dissolve the copper sulfate separately in about 100 ml of water and your the solution slowly into the other liquid with containst stirring. Add the ferrocy and solution cool and didute to exactly 1 liner Of the various constituents the copper salt only need be weighted with exactness. Twenty five ml of the respent are reduced by 30 mg of glucose.

CALCULATION The 25 ml of copper solution is reduced by exactly 50 mg of glucose Therefore the volume run out of the burct to effect the reduction contained 50 mg. of the sugar. The formula for calculating the percentage of the sugar is the following $\frac{0.000}{1000} \times D \times 100$ = per cent in original sample wherein x is the number of ml

of the diluted urine required to reduce 2s ml of the copper solution and D is the dilution of the urine (D equals 1 for undiluted urine 10 for urine diluted i 10 etc)

In the use of this method chloroform must not be present during the titration, if used as a preservative in the urine it may be removed by hoiling a sample for a few minutes and then diluting to its original volume

Interpretation. Glucose in the unne in amounts sufficient to be detected by the commonly employed qualitative tests (i.e., 01 to 02 per cent or more) ordinarily indicates a pathological condition, although it must be remembered that benign glucosuma is not uncommon (see p 823 for further discussion), and that other reducing sugars (lactose, pentose) cannot be distinguished from glucose by the ordinary reduction tests, either qualitative or quantitative Persistent glucosuria may indicate diabetes mellitus, a disorder in which the amount of sugar may rise as high as 10 per cent and averages 3 to 5 per cent. The volume of unne excreted per day is usually also large and the absolute sugar excretion may thus be very great (100 g of glucose per day are not uncommon) The quantitative methods for the estimation of sugar in unne enable us to determine the severity of this disorder as well as to follow its course under treatment, etc

2 Sumner's Method. 127 Principle Unne is heated with a dimitrosalicy lic acid reagent which is reduced by the sugar and the resultant color is compared with standards This is a rapid method, applicable to normal as well as gly cosure urne but is not as precise as the more elaborate method of Shaffer and Hartmann described below

Procedure Pipet into a Folin-Wu blood sugar tube 1 ml of urine (diluted if necessary) and 3 ml of the dinltrosalicytic acid reagent in Mix and heat 5 minutes in boiling water Cool 3 minutes in running water, dilute to 25 ml mix and compare in a colorimeter with a standard glucose solution treated simultaneously in the same way A concentration of 0 1 per cent glucose is satisfactory for the standard solution If the color obtained with the urine is too dark, repeat the test using more dilute urine

CALCULATION

 $\frac{\text{Reading of Standard}}{\text{Reading of Luknown}} \times \frac{\text{Dilution}}{10} = \text{per cent glucose}$

3 Folin McEllroy Peck Method Principle The method is a titration procedure depending upon the use of an alkaline copper solution in which the cupric

¹³⁷ Sumner J Biol Chem 45 383 (192a)

¹¹⁶ To 10 g. crystallized phenol add 22 ml 10 per cent \sOII Dissolve in a little water and dilute to 100 ml Weigh 69 g. sodium basilite and to this add 69 ml of alkaline phenol solution. To this add a solution containing 300 ml 40 per cent \sold 200 k Rochelle salt (\sKC.H.O. 4H.O) and 850 ml t per cent dinitrosales lie acid Mix and keep tightly stoppered in well filled bottles. The reagent should keep for at leat one year Dinitrosalicylic acid may be obtained from the Eastman Kodak Co Rochester

hydroxide is held in solution by means of phosphate instead of the customary tartrates, citrates, or glycerol. The method is applicable to the determination of factose in milk. 129

4 Method of Shaffer and Hartmann ¹⁴⁰ Principle The sugar solution is boiled with Fehling's alkaline copper solution (as modified by Sokhlet) under the conditions prescribed in the standard method of Vlunson and Whiker whose tables may therefore be used in the calculation of results. The residual cupric salt may then be converted into cuprous iodide with the bheration of an equivalent amount of iodine $2Cu^{++} + 4I^{-} + 2Cu^{+} + 1$. Of the cuprous salt may be oxdized in the presence of a known amount of iodine $2Cu^{+} + 1_1 \rightarrow 2Cu^{++} + 2I^{-}$ Iodine liberated, or excess iodine found in the second case may be ittrated with sodium thosulfate. In the titration of cuprous copper, oxalate is added to depress the ionization of the cupric salt. Shaffer and Hartmaon prefer the cuprous titration

Procedure. Pipet 25 ml of each of the two Fehinog's solutions. Into a 300-or 400-mi flask. Add 50 ml or less of the (approximately neutral) sugar solution containing from 20 to 200 mg. of sugar, and water (if necessary) to make a total volume of 100 ml. Cover with a small inverted beaker and heat on an asbestos matover a flame so adjusted as to bring the solution to boiling in 4 minutes. Boil 2 minutes. Stand the flask in the sink under running water till cool (3 to 4 minutes). Then use one of the two following procedures.

Guprous Titration Add 50 ml (accurate pipet) or 25 ml If but little cuprous oxide is present, of iodate-iodide solution, in followed by 15 to 17 ml of 5 N H₂SO. The acid should be added from a cylinder or fast-flowing pipet in order that acidification of the whole solution may be accomplished promptly Shake the solution gently for a few moments until the cuprous oxide has dissolved. The solution should become clear, but some cuprous lodide may separate Add 20 ml of saturated solution of potassium oxilate and rotate the flask until the cuprous lodide is completely dissolved.

Titrate with 0 1 N sodium thiosuifate (see Appendix) adding a few mi of starch solution toward the end before the disappearance of the green color Run a blank by holling the Fehling's solution with 50 ml. of water instead of sucar solution

Calculation From the blank titration subtract the titration of the sugar determination, the remainder representing 1- required for the oxidation of the cuproussalt Multiply by the copper factor of the thosulfate (I ml of 0.1 N = 6.38 mg of Cu), and find the amount of sugar equivalent to the copper by reference to Munson Walker tables 14 or divide the amount of copper reduced by the corresponding ratio obtained from the glucose curve in Fig. 252

Cupric Titration. To the choled alkaline copper solution add 6 g of KI and 25 mi, of 5 N H₂SO₄. Titrate with standard thiosulfate, adding starch

¹³³ Folin and Peck J Biol Chem. 38, 287 (1919)

¹⁴⁵ Shaffer and Hartmann J Biol Chem 45 365 (1921)

¹⁴¹ Fehing s Solutions (Sozhlet Modification) (I) Dissolve 34 64 g CuSO, 511,0 in water to make 500 ml (II) Dissolve 173 g Rochelle salt and 50 g NaOll in water to make 500 ml

¹⁴² Iodate-Iodide Sol tion. Dissolve 5.4 g. klOs and 60 g. kl in water to which a small amount of alkali has been added and dilute to a liter

¹⁴² Methods of Analysis of the Association of Official Agricultival Chemists 7th ed. Washington D.C. 1950.

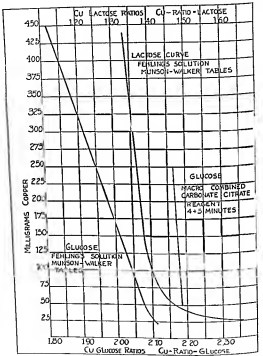


FIG 252 COPPER GLICOSE RATIOS FOR FEHLING'S SOLUTION AND FOR THE MACRO COMBINED REAGENT AND COPPER LACTOSE RATIOS FOR FEHLING'S SOLUTION.

Divide the amount of reduced copper by its corresponding ratio as shown by the respective curves

solution toward the end. The titration is subtracted from a similar blank determination on the Fehling's solution, the difference representing copper reduced by the sugar. For the cupric titration the copper solution must be measured accurately. Consult tables or chart (Fig. 252) for sugar values.

5. Hones' Modification of the Hogedorn-Jensen Method: Principle. The Hagedorn-Jensen method (see Chapter 23) is modified so that it may be used for larger amounts of glucose (1 to 3 mg) Maltose may also be determined in the presence of starch Power and Wilder adapt the method to the determination of 1 to 50 mg of glucose in urine. For accurate results the urine must be cleared with increury. For an accuracy within 1 to 5 per cent on diabetic urines the uncleared urine may be used, a correction of 0 25 per cent glucose being subtracted.

Procedure. Into a test tube (1×7 in.) pipet 5 ml. of Solution A (alkaline ferricyanide). Add 5 ml. of unknown solution (if necessary add water to make the 5 ml. volume). In another tube put 5 ml. of water and 5 ml. of Solution A. Incline the tube to mix in any drops of Ilquid adhering to the sides. Cover with glass bulbs (with about an Inch of tubing left attached). Place for 15 minutes in a boiling water bath 2 to 3 in. deep. Cool for 3 minutes in cold running water.

Add 5 ml, of Solution B and then 3 ml, of Solution C, using rapid pipets which need not be precise. Run in thiosulfate to a pale vellow color, add a few drops of starch, and titrate to disappearance of the blue color, Subtract titration of unknown from titration of the blank. Convert result into mi, of 0.01 N thiosulfate, Consult Fig. 253 for amounts of glucose or maltose. In the presence of starch stir more vigorously and titrato more slowly. The maltose values are also easily calculated by multiplying the number of ml. of 0.01 N thiosulfate used by 0.414 mg. invert sugar and thus indirectly sucrose may also be determined. The factor for invert sugar using a final volume of 15 Instead of the usual 10 ml, is 0.347 and for sucrose 0.329. For a standard invert sugar solution dissolve 0.95 g. of sucrose in 150 ml. of water, add 30 ml. of 0.5 N HCl, heat to boiling, and boil 1 minute. Cool rapidly, add 30 ml, of 0.5 N NaOH, and dilute to 500 ml, 100 ml, will contain 0.2 g, of invert sugar. Other sugars may also be determined. The factor for fructose is 0.32, for ribose or an equimolecular mixture of glucose and galactose 0.38, arabinose 0.35, mannose and rhamnose 0.34, xylose 0.33, lactose and galactose 0.43.

6. Benedict's Picrote Method (Adopteo by the Committee ov Urinary In-Pariments of the Association of Life Insurance Madical Directors of America). Principle. The color produced by the reduction of piene acid (p 67) is compared with permanent morganic standards representing definite concentrations of sugar.

¹⁴⁴ Solution A. Dissolve S. 25 g. potassum ferrey ande and 10 0 g. auhvdrous sodium carbonate in water to make 1000 ml. Store in an epoque bottle and keep 2 to 3 days before use Solution B. Dissolve 1.5 g. Kl. 25 0 g. ane sulfate, and 125 g. NoCl in water to make 500 ml. Traces of codine appear in the solution on stering Remove be filtering through 2 thicknesses of filter paper. Solution C. Didute 5 ml et glacula actic and to 100 ml. Starch Str up 1 g. of Merck solutible starch with 20 ml water, add to 50 ml boding water, boil 2 minutes, cool, and make up to 100 ml. Will keep for several months Sodium Thoculfate in approximately, N/75 solution weed m a 10-nl buret graduated in 0.02-ml divisions. Dissolve 3 33 g of the salt in 10 liters of boiled-out water Keep in bottle protected by sodalime tube and run into buret by siphen Standardine seach day at first, then every three to four days. Pipet 5 ml 0.02 N KlO₃ solution (715 g in 1000 ml.) into a tube Add 5 ml. 2 recent kill and 3 ml. 5 per cent acetie and Tittate busing starch as indicator.

Procedure Measure 1 ml of urine into a test tube graduated at 25 ml Ad-3 ml of picric acid solution (2 g of pure dry picric acid per liter) and 0 5 m of 5 per cent NaOH Add next 5 drops of 50 per cent acetone solution (pre pared fresh each day by diluting acetone with an equal volume of water and place the tube promptly in a boiling water bath. In 12 minutes remov

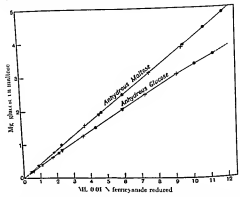


FIG 203 CHART FOR SEGAR VALUES BY HANES METHOD

the tube, cool, and dilute the contents to 20 ml Compare this colored solu tion with the permanent standards in tubes of the same dimensions, and estimate the amount of sugar The permanent sugar standards are made 25 described below

PERMANENT SUGAR STANDARDS145

Put each standard in test tubes of the same diameter as used in sugar de termination

Dissolve 200 g. ferric chloride (FeCl; GHrO) in about 300 ml. distilled water transfer to 500-ml volumetric flack, make up to volume with distilled water and mix well. Filter through a dry filter paper

Cobalt Chloride (analyzed grade)

Dissolve 1.0 g. cobalt chloride (CoClr 6HrO) in about 300 ml distilled water transfer to a .00-ml volumetric flack, make up to volume with distilled water and mix well-Filter through a dry filter paper

Dilute Hydrochloric Acid

Dilute 5 ml. concentrated by drochlorse and to 30 ml with distilled water

¹⁴³ Solutions needed.

Ferric Chloride (Merck & Analyzed)

Sugar %	Ferric Chloride Solution ml	Cobalt Chloride Solution ml	Dilute Hydrochloric Acid ml	Water	
0 1	18	7	S		
0 2	28	13	8		
0 3	22	22	8	to make 100 ml	
0 4	16	30	8		
0 5	14	40	8		

7. Hawkins and Van Slyke Method.¹⁴⁴ Principle The time required for the urine to decolorise potassium ferricynnide solution is an index of the amount of reducing sugar present Since normal constituents in concentrated urine may give reduction equivalent to as high as 0.4 per cent glucose, the method is regulated to determine reducing substances in concentrated urines in concentrations of 0.5 per cent and above, and in dulter time in concentrations of 0.5 per cent and above.

Procedure. Dilute 1 ml. of urine to 50 ml. If urline is known to be high in sugar (over 3 per cent) dilute 1 100, if low in sugar, dilute 1 25. Albumin need not be removed Pipet 2 ml. of diluted urine into n pyrex test tube (14 × 125 mm outside dinmeter). Add 2 ml of ferricyanide solution. Mix. Immerse in a beaker of boiling water. A similar test tube containing water is immersed for comparison. Make a white background by pasting paper on the side of the beaker away from the observer. A casserole may be used instead of n beaker. Determine the time in seconds required for the last trace of yellow disappear, preferably using a stop watch. From the chart, Fig. 254, obtain the percentage of sugar in the urine. The chart is for a dilution of 1 50. If a 1 25 dilution is used, divide the result obtained from the chart by 2; for a 1 100 dilution, multiply the result by 2.

8. Fermentation Method: Principle. This method consists in the measurement of the volume of carbon dioxide evolved when the dextrose of the urine undergoes fermentation with yeast None of the various methods whose manipulation is based upon this principle is absolutely accurate The method in which Linhorn's saccharometer (Fig. 15, p. 69) is employed is perhaps as satisfactory as any for chincal purposes

Procedure. To about 15 ml of urine in a mortar, add about 0.25 g. of active dry yeast and mix thoroughly. Transfer the mixture to the saccharometer,

iii Ilankins and Van Slyke J Biol Chem 81, 459 (1929) For a similar procedure using smaller volumes of solution (and therefore with a slightly different calibration curve) which is also directly applicable to tungsite a

J Biol Chem. 129, 51 (1937)
in Potanum Ferrepannic Raggent Dissolve 75 g of anhydrous potassium carbonate in the potanum Ferrepannic Raggent Dissolve 150 ml of water Dissolve 1 g of potassium the properties of the properties of the potantial state of the potantial state of the properties of the prop

being careful that the graduated tube is completely filled and that no air bubbles gather at the top. Allow the apparatus to stand in a warm place 30° C) for 12 hours, and observe the percentage of dextrose as indicated by the graduated scale of the instrument Both the percentage of dextrose and

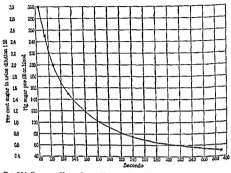


Fig 254 Chart of Urine Sugar Values for Hawkins-Van Slike Method J Bool Chem 81 459 (1929)

the number of cubic centimeters of carbon dioride are indicated by the graduations on the side of the saccharometer tube. Controls should be run using normal urine and such urine to which sugar has been added

9 Polariscopic Examination Before subjecting urine to a polariscopic examination the slightly acid fluid should be decolorized as thoroughly as possible by the addition of a little base lead acetate. The urine should be well stirred and then filtered through a filter paper which has not been previously moistened. In this way a perfectly clear and almost colorless liquid is obtained.

In determining dextrose by means of the polariscope it should be borne in find that this carbohydrate is often accompanied by other optically active substances such as proteins fructose \$\tilde{\theta}\)-hydroxybutyric acid, and conjugate glucuromates which may introduce an error into the

Specific Rotation f.barnee + 52 49 Maltose +136 5 1somaltose + 68 0 Lactore + 52 53 Fructose - 92 25 ≠llydroxybutyric acid ~ 24 12 Conjugate i glucuronic Leverotatory in acida varying degrees polariscopic reading, the method is however, sufficiently accurate for practical purposes

For directions as to the manipulation of the polariscope, see p 69.

The table gives the specific retations of some physiologically im-

tations of some physiologically important sugars as well as of certain other optically active substances the possible presence of which must be borne in mind in determining glucose polarimetrically in urine

DETERMINATION OF SUGAR IN NORMAL HRINE

Principle Since the nature of normal urine sugar (i.e. reducing substances) is not definitely established at is not possible to state which of the methods of determina tion is to be preferred. The methods involve removal of interfering substances and colorimetric determination of reducing nower

Procedure

precipitation.

(a) REMOVAL OF INTERFERING SUBSTANCES METHOD OF FOLIN AND SVEDBERG 141 To 5 ml of urine in a 50 ml Erlenmeyer flash add 5 ml of 0 05 N oxalic acid, 10 ml of water, and (last) 1 5 g of Lloyd s alkaloidal reagent 100 Shake gently for 4 minutes. Filter through a quantitative paper into another small flash containing 2 g of Permutit Shake 3 minutes Decant

(b) DETERMINATION OF SUGAR. Sugar may be determined in filtrates from treatment with Lloyd's reagent by one of the methods used for blood sugar 150 (see Chapter 23) The method of Sumner (see p 920) is said to be applicable to urine without previous treatment. The urine may also be fermented and fermentable sugar determined by measurement of CO1 given off or by differ ence To 10 ml of the filtrate from treatment with Lloyd's reasent add 3 ml of a suspension of 1 cake of compressed yeast in 20 ml of water Keep at 37° C for 40 minutes Filter and determine sugar Fermentable sugar is usually about 0 01 per cent (see Chapter 28) The urine may also be hydro lyzed and total sugar determined To 8 ml of urme filtrate add 1 ml of 2 6 N H2SO, and heat in a boiling water bath for 75 minutes Add 1 ml of silica free NaOH exactly equivalent to the acid added Determine sugar by any of ahove methode

PROTEINISI

1 Colorimetric Determination of Proteins (Method of Hiller, McIntosh, ond Van Slyke) 181 Principle The protein is precipitated with trichloroacetic acid 181 dissolved in NaOH solution, and copper ions added to give a hieret color which is compared with a standard

Procedure Measure 2 ml of the urine (previously adjusted to a pH of about 7 4) into a centrifuge tube, add an equal volume of 10 per cent triciloroacetic acid solution, mix and centrifuge for 5 minutes If the volume of the pre cipitate is less than 0 2 or more than 0 6 mi , repeat using more or less urine Pour off the supernatant fluid Dissolve the precipitate in about 3 mi of 3 per cent NaOH and wash into a tube graduated at 10 ml with portions of the 3 per cent NaOH until the volume is about 9 mi Add 0 25 mi of 20 per cent

¹⁴⁵ Folin and Svedberg J Biol Chem 70 405 (1926) Hamilton (J Biol Chem 78 63 (1928)) onuts Permutit but uses Lloyd's reagent extracted with concentrated HCl for

¹ day was de with water and the extracted with concentrated Who, for or of day
1 doy lost of with water and the extracted with concentrated Who, for or of day
1 doy lost of the with water and the extracted with concentrated Who, for or of day
1 doy lost of the with water and the with water of the with See bol of J Bud Chem 68 706 (1926)
1 The method for album 1 adopted by the Committee on Urman 1 Impairments of the

Association of Life Insurance Medical Directors of Musers as given on p 1-9
Hiller McIntosh and Van Siske J Clin. Intest 4 235 (19-7)
Hiller McIntosh and Van Siske J Clin. Intest 4 235 (19-7) (1343)) from 4 to 20 per cent of the protein in prote numa may not be precipitable by friel loroacetic ac d. This is a account for discrepance six protein determination between methods involving irichloroscetic send precipitation and those not involving such

copper sulfate solution and dilute to 10 ml with the \aOII \text{ \text{ Nix by shaking.}} Let stand 10 minutes Centrifuge and compare with a standard For a stand ard measure 5 ml of a biuret solution containing 13 33 mg of biuret, equivalent to 12 3 mg of protein, into a tube graduated at 10 ml \text{ \text{ Add distilled water}} to 8 ml, 1 ml of 30 per cent \aOII, 0 25 ml of copper sulfate, and water to make 10 ml \text{ \text{ III}}, let stand for 10 minutes, centrifuge, and compare la secolorimeter with the unknown, setting this standard at 15 mm

For determining albumin add to 10 ml of the urine, at pil 7 4, 10 ml of 44 per cent sodium sullate solution, mix well, and put in an incubator at 37°C for three hours Filter Proceed as for total protein, using four times the volume The dlobulin is estimated by different by

CALCULATION For total protein

$$\frac{1a}{R} \times \frac{123}{m_1 \text{ prime post}} = g$$
 protein per liter

For albumin

$$\frac{\text{La}}{R} \times \frac{123 \times 2}{\text{ml. pitrate need}} = \text{g. albumin per later}$$

2 Folin's Gravimetric Method for the Determination of Protein in Unit-Principle. The protein precipitated by heat and actic acid is centrifuged washed, and wighed.

Procedure Pipet 10 ml of urine into an ordinary conical centrifuge tube which has been previously weighed add 1 ml of 5 per cent acetic acid, and let stand for 15 minutes in a beaker of boiling water At the end of this mar remove the tube from the water bath, and centrifuge for a few minutes. Pour off the supernatant liquid, stir up the precipitate in the tube with about 10 ml of boiling 0 5 per cent acetic acid, and again centrifuge. Remove the supernatant liquid from the precipitate in the tube and wash once more this time with 50 per cent alcohol. After centrifuging, pour off the supernatant alcohol and place the tube for 2 hours in an air bath at 100 to 110° C.

The precipitate may also be filtered off and washed on a small tared filter paper, then dried and weighed in this case it is better to use a larger volume of urine. Instead of weighing, the nitrogen in the precipitate may be determined by the Kieldahl method 160.

CALCULATON Multiply the we glot of the precipitate in grams by 10 to obtain the percentage of protein present If a volume other than 10 ml, of urine is used multiply by the appropriate volume factor

Interpretation The amount of albumin occurring in the urine is not necessarily an index of the eventh or type of the disorder giving rise to it. Hence no significant figures can be given Normal human urine probably contains a trace of albumin which is too slight to be detected or determined by the ulual procedures. The determination of albumin may be of assistance in following the course of kidney disturbances but the results can be interpreted only in the light of other clinical findings. (See discussion under "albumin, p. 828.)

¹⁴ In order to arrive at correct figures for the protein content it is then only necessary multiply the total nitrogen content by 6.2.5 (see p. 239) Correction should be made for the nitrogen content of the filter paper used unless this factor in negligible.

 Esbach's Method: Principle. This method depends upon the precipitation of principle and measurement of its volume after settling 'The apparatus used is Esbach's albuminometer (Fig. 255)

Procedure. In making a determination fill the albuminometer to the point U with urine, then introduce Eshach's reagent 138 (or 10 per cent trichloro-

acetic acid, as recommended by Qulck) until the point R is reached. Now stopper the tube, invert it slowly several times in order to insure the thorough mixing of the fluids, and stand the tube aside for 24 hours. Creatinine, resin, acids, etc. are precipitated in this method, and for this and other reasons it is not so accurate as the coagulation method. It is, however, extensively used clinically. According to Sabil, Esbach's method is "accurate approximately to 1 part per 1000," whereas Pfeifler claims it is not accurate for less than one-half or for more than 5 parts per 1000.

CALCULATION The graduations indicate g of protein per liter of urine. Thus a reading of 3 indicates 3 g per liter, or 0.3 per cent protein.

4. The Determination of Albumin (Method Adopted by the Countries of Uninary Impairments of the Association of Life Isburance Medical Dibectors of America). Principle. Clarified urine is treated with sulfosalicyhe acid and the degree of turbidity produced is measured by comparison with artificial standards.



Fio 255 Essacu's ALBUMINOMETER.

Procedure. Pipet 2.5 ml, of centrifuged urine into a test tube graduated at 10 ml. and add 3 per cent sulfosalicylic acid (30 g. in 1000 ml, of distilled water) to the 10-ml. mark. Invert the tube to mix, allow to stand 10 minutes, and compare the turbidity with the permanent turbidity standards. Record the value of the standard most closely matched, as the albumin content of the urine.

Preparation of Permanent Albumin Standards. 114 Dissolve 20 g. of gelatin (Super X brand in sheet form, Coignet Chemical Products Co., New York) in 120 to 140 ml. of distilled water at 45° to 55° C. and make up to 200 ml. Add about half the white of an egg and stir in. Heat in a water bath for at least 30 minutes after a temperature of 90° C. has been attained. Filter hot through a Whatmani's No. 4 paper yielding a perfectly clear, slightly yellow solution. Immediately before using, add 0.3 ml. of formalin (40 per cent formaldehyde solution) to each 100 ml. of gelatin solution. Formazin, the material to be suspended in the gelatin, is made as follows. Dissolve 2.5 g. of urotropin in 25 ml. of distilled water at room temperature. Add to 25 ml. of 1 per cent hydrazine sulfate solution also at room temperature. Mis, stopper, and allow to stand at least 15 hours. Suspend the white amorphous precipitate uniformly by gently inverting the flash a few times. Add 14.5 ml. of the formazin suspension to 100 ml. of 10 per cent gelatin solution at

¹⁴⁴ See Appendix

¹⁸ These permanent standards were developed in the laboratories of the Metropolitan Ide Insurance Company and the Mutuel Benefit Ide Insurance Company and detailed description of the preparation of these standards, see Kingabury, Clark, Williams, and Post J Lab Clin Med. 11, 981 (1920).

i5° to 55° C. (to which the correct amount of formalin has been added) and mix thoroughly. This produces a turbidity equivalent to that made by an albumin solution of 0.1 per cent, or one containing 100 mg. of albumin in 100 ml. when precipitated by 3 volumes of 3 per cent sulfosalicylic acid. Dilute this stock suspension as follows to make the remaining standards

Stock Formazin Suspension Equivalent to 100 mg Albumin per 100 ml	10% Clarified Gelatin	Value of Standard Made	
ml	ml	%	mg
25 0	26	0 05	50
20 0	30	0 04	40
15 0	35	0 03	30
10 0	40	0 02	20
5 0	45	0 01	10
2 5	55	0 005	5

Pour each standard into a teat tube of the same dimensions as those used in making the test with urme. Seal the tube with a waxed stopper and allow to cool to room temperature. In a short time the gelatin should solidify and after a few days cannot be melted at any room temperature, in extremely hot weather put the tubes in a cool place for a few days before any attempt is made to use them. Keep the standards in a well-lighted room if in time they become greenlsh, they may be bleached by exposure to sumlight without changing their turbidimetric value. The standards may be checked against sheep serum standards of known protein content precipitated in the same manner as in the urine test. Standards older than 8 months should be replaced unless actual tests show that they have their original degree of turbidity. In the course of a year there may be only a very slight diminution in the turbidimetric value of the standards. There is no detectable change in 6 to 8 months.

ACETONE BODIES

1. Van Slyke's Methods. ** Principle The method is based on a combination of Shaffer's oxidation of shydrorybutyne and to acetone (p. 934), and Denge's precipitation of acctone as a base mercure suifate compound Glucose and certain other interfering substances are removed by precipitation with copper sulfate and calcium hydroxide Preservatives other than toluene or copper sulfate should not be used.

calcium hydroxide suspension, shake, and test with litmus. If not alkaline, add more calcium hydroxide. Dilute in the mark and let stand at least one balf hour for glucose to precipitate. Diter through a dry foided filter. This procedure will remove up to 8 per cent if glucose. Urine containing more should be diluted enough to bring the glucose down to 8 per cent. The copper treatment is depended upon to remove interfering substances other than glucose, and should therefore never be omitted, even when glucose is absent. The filtrate may be tested for glucose by boiling a little in a test tube. A precipitate of yellow cuprous oxide will be obtained if the removal bas not been complete. A slight precipitate of white calcium saits always forms, but does not interfere with the detection of the yellow cuprous oxide.

Determination of Total Acetone Bodies (Acetone, Acetoacetic Acid, and 8-Hydroxybutyric Acid). Place in a 500-ml. Erlenmeyer flash 25 ml. of urine filtrate, Add 100 ml, of water, 10 ml. of 50 per cent sulfuric acid, and 35 ml. of the 10 per cent mercuric sulfate. Or, in place of adding the water and reagents separately, add 145 ml. of the combined reagents Connect the flash with a reflux condenser baying a straight condensing tube of 8 or 10 mm diameter, and beat to boiling. After boiling has begun, add 5 mi, of the 5 per cent bichromate through the condenser tube. Continue boiling gently 11/2 hours. The yellow precipitate which forms consists of the mercury sulfate-obromate compound 150 of acetone (total), it is collected in a Gooch or "medium density" alundum or fritted glass crucible, washed with 200 ml of cold water, and dried for an hour at 110° C. The crucible is allowed to cool in room air (a desiccator is unnecessary and undesirable) and weighed Several precipitates may be collected, one above the other, without cleaning the crucible. As an alternative to weighing, the precipitate may be dissolved and titrated as described below.

Determination of Acetone and Acetonacetic Acid. The acetone plus the acetoacetic acid, which completely decomposes into acetone and CO_1 on heating, is determined without the β -bydroxybutyric acid exactly as the total acetone bodies, except that (i) no bichromate is added to oxidize the β -bydroxybutyric acid and (2) the boiling must continue for not less than 30 nor more than 45 minutes Boiling for more than 45 minutes splits off a little acetone from β -bydroxybutyric acid even in the absence of chromic acid, in

Determination of β -Hydroxybutyric Acid. The β -hydroxybutyric acid alone is determined exactly as total acetime bodies except that the preformed acetione and that from the acetoacetic acid are first boiled off To do this

its This contains about 77 per cent mercury and in the absence of chromate has approximately one of the following formulas 3HgSO, 5HgO 2(CH₂)₂CO or 2HgSO, 3HgO (CH₂)₂-

Is Blank Determination of Precipitate from Substances in Urine Other than the Acctome Boiles The 25 ml shquot of unine filtrate is treated with sulfurire and and water and boiled 10 minutes to drive off acctome The residue is made up to 175 ml with the same amounts of mercuric sulfate and sulfure and used in the above determinations, but without chromate and is build under the reflux for 45 minutes Longer boiling splits off some acctone from \$h_i\text{drox}\text{builty inc} \text{acid}\text{, and must therefore be avoided The weight of Precipitate obtained may be subtracted from that obtained in the above determination

The blank is so small that it appears to be significant only when compared with the small amounts of acctone bothes found in normal or nearly normal urines. In routine analyses of diabetic urines it is not determined

Tests of Reagents: When the complete total acctone bodies determination including the preliminary copper sulfate treatment is performed on a sample of the similar distribution of the preliminary copper sulfate treatment is performed on a sample of the builded water with the obtained. This test must not be mutted

the 25 ml. of urine filtrate plus 100 ml. of water are treated with 2 ml. of the 50 per cent sulfurle acid and boiled in the open flask for 10 minutes. The volume of solution left in the flask is measured in a cylinder. The solution is returned to the flask, and the cylinder washed with enough water to replace that boiled off and restore the volume of the solution to 127 ml. The 8 ml. of the 50 per cent sulfuric acid and 35 ml of mercuric sulfate are added. The flask is connected under the condenser and the determination is continued as described for total acctone bodies.

Titration of the Precipitate in the Abose Methods. Instead of weighing the precipitate, one may wash the contents of the Gooch, including the asbestos, into a small beaker with as little water as possible, and add 15 ml. of normal IICI. The mixture is then beated, and the precipitate quickly dissolves in case an alundum or glass crucible is used, it is set into the beaker of acid until the precipitate dissolves, and then washed with suction, the washings being added to the beaker. In place of using either a Gooch or alundum crucible, one may, when titration is employed, wash the precipitate without suction on a small quantitative filter paper, which is transferred with the precipitate to the beaker and broken up with a rod in 15 ml of normal IICI

In order to obtain a good end point in the subsequent titration it is necessary to reduce the acidity of the solution. For this purpose it has been found that the addition of excess sodium acetate is the most satisfactory means Six to 7 ml. of 3 M acetate are added to the cooled solution of redissolved precipitate. Then the 0 2 \1 KI is run in rapidly from a buret with constant stirring If more than a small amount of mercury is present, a red precipitate of light at once forms, and redissolves as soon as 2 or 3 ml. of Ki in excess of the amount required to form the soluble Killgl, has been added. If only few mg of mercury are present, the excess of Kl may be added before the ligi, bas had time to precipitate so that the titrated solution remains clear In this case not less than 5 ml of the 0 2 M Kl are added, since it has heen found that the final titration is not satisfactory if less is present. The excess of Kl is titrated back by adding 0 05 M ligCl: from another buret until permanent red precipitate forms Since the reaction involved is 11gCls + 4Kl - Kilgl. + 2KCl, 1 ml of 0 05 M ligCl. is equivalent in the titration to 1 ml. of the 9.2 M Kl.

In preparing the two standard solutions the 0 05 M ligCl₂ is standardized by the sulfide method, and the lodine is standardized by ttration against it A slight error appears to be introduced if the lodide solution is gravinetric cally standardized and used for checking the mercury solution, instead of vice versa.

In standardizing the mercuric chloride the following procedure bas been found convenient 25 ml of 0 65 M HgCls is measured with a calibrated pipel; diluted to about 100 ml, and His B is run in until the black precipitate floculates and leaves a clear solution. The HgS, collected in a Goodb crucible and dried at 10°C, should weigh 0 2982 of the solution is accurate

Both by gravimetric analyses of the basic mercuric sulfate acetome precipitate and by titration, the mercury content of the precipitate has been found to average 76 pper cent. On this basis, each mi, of 9 2 M KI solution, being equivalent to 10 0 mg of Hg, is equivalent to 13 0 mg of the mercury acctone precipitate

Titration is not quite so accurate as weighing, but, except when the amounts determined are very small, the titration is satisfactory.

CALCULATION One mg. of \$\beta\$-hydroxybutyra scall yields \$45 mg of precupitate One mg. of acctone yields 20 0 mg. of precupitate One ml of 0 2 M M solution is equivalent to 13 mg. of precupitate in titration of the latter

In order to calculate the nectone bodies as β hydroxybutyric acid rather than acctone use the above factors multiplied by the ratio of the molecular weights $\frac{\beta}{8}$ acid $\frac{104}{88} = 1.793$ In order to calculate the acctone bodies in terms of molecular concentration divide the factors in the table by 88. To calculate ml of 0.1 M acctone bodies per liter of uring use the above factors multiplied by 10 000/88 or 172.4

SPECIAL FACTORS FOR CALCULATION OF RESULTS WHEN 25 ML OF URINE FILTRATE EQUIVALENT TO 2.5 ML OF URINE ARE USED FOR THE DETERMINATION

Determination Performed	Acetone Bodies Calculated as g of Acetone per Liter of Urine Indicated by		
	I g of precipitate	1 ml of 02 W KI sol	
Total acetone bodies* β Hydroxybutyric acid \text{\text{cetone}} + acetoacetic acid	21 8 26 4 20 0	0 322 0 344 0 260	

^{*}The total acetone hodies factor is calculated on the assumption that the molecular proportion of them in the form β h, droxy buty he acid is $^{-}$ 0 per cent of the total which proportion is usually approximated in acetone needs β h, droxy buty it is each yields only 0.75 molecule of acetone the factors are strictly accurate only when this proportion is present but the error introduced by the use of the approximate factors is for ordinar purposes not serious. The actual errors in percentage of the amounts determined are as follows molecular proportions of acetone hodes as β and 0.50 error 6.5 per cent β acid 0.90 error 1.3 per cent β acid 0.90 error 1.3 per cent.

Interpretation. Normal adults on a mixed diet excrete on the average 3 to 15 mg of combined acetone and acetoacetic acid per day, and any thing over 20 mg is usually pathological Usually about one fourth of this total is acetone, although the proportion varies considerably. The amount is considerably increased in fasting and on a carbohy drate-free diet due to the development of ketosis. In severe diabetic ketosis values up to 6 g per day or even higher may be noted. It is sometimes found in large amounts in intovications associated with pregnancy. It may be found in increased amounts in the urine in a great variety of pathological conditions. Quantitative estimation enables us to follow the course of the ketosis. Ammonia excretion is also largely increased in these conditions, being used in the neutralization of the excess acids formed in the body. (See also Chapter 29.)

β Hydroxybutyme acid may occur in normal human urine to the extent of 20 to 30 mg per day In fasting or on a carbohydrate-free diet, very large amounts may be exercted (up to 20 g per day) In severe diabetes mellitus the largest amounts are found, and exerctions of 50 or even 100 g or over per day have been noted In this condition it is usually the most abundant of the acetone bodies making up from 60 to 80 per cent of the total. The ratio is, however, by no means constant, and it should be borne in mind that in rare cases large amounts of β-hydroxy butyme acid may be eliminated although the acetone exerction is very low. It is always present in the urine when large amounts of acetone are present

A discussion of disturbance of acid base equilibrium due to abnormal

f it metabolism may be found on pp 692 and 693

2 Method of Shaffer and Marriott 144 Principle By this procedure the com bined acctone and acctoacctic acid is determined in the same sample of urine used in the determinat on of \$\beta\$-hydroxybutyric acid. The preformed acctone and the acctoacetic ac d are distilled off together as acetone an I determined by the jodine titration method It c s hydroxybutyrie acid remains in the resilie from distillation and is oxidized by means of potassium bichromate. The product of the oxidation is acetone which is distilled off and determined as such

Procedure Determination of Acetone and Acetoacetic Acid Measure from 25 to 100 ml 112 or more of urine (usually 50 ml) with a pipet into a 500 ml volumetric flask containing 200 to 300 ml of water Add basic lead acetate solution (USP) in volume equal to that of the urine used113 and mix well Add concentrated ammonium hydroxide in amount equal to about one half that of the lead acetate solution Dliute the contents of the flask to the mark with water, shake and let stand for a few minutes. Then filter the liquid, preferably through a folded filter Measure 200 mi of the filtrate into a round bottomed flack (800 ml or liter Kjeldahl flasks are consenient), dliute with water to about 600 ml , and add 15 ml of concentrated sulfuric acid and a little tale or a boiling stone Distil until about 200 ml of distillate have been collected The tube of the condenser should dlp beneath the surface of the water in the receiving flask so that no loss of acetone will occur The distilling flask must also be fitted with a dropping tube or drop ping funnel so water may be run in from time to time and the volume of liquid in the flask kept from becoming less than 400 to 500 ml A good condenser should be used, but it is not necessary to cool the distillate in ice

The distillate thus obtained is transferred to a second Kjeldahi flask and 10 ml of 10 per cent NaOH added It is then redistilled for about 20 min utes 144 The second distillate is then titrated with standard lodine and thio sulfate solutions

This is done by adding 10 to 25 ml of 0 1 \ lodine solution (0 02 \ if the amount of acetone bodies as indicated by a qualitative test is small) and 10 mi of strong NaOil (about 40 per cent) Let stand for 10 minutes Add 18 ml of concentrated HCl Titrate with @ I N sodium thiosulfate solution to a pale yellow color, add a few ml of soluble starch solution (see Appendix) and con tinue titration to disappearance of the blue color

CALCULATION Subtract the number of ml. of 0.1 \ thiosulfate solution used from the volume of 0 1 \ iodine solution employed Since 1 ml of the iodine solution 13 equivalent to 0 967 mg of acctone and since 1 mi of the thiosulfate solution is equiva lent to 1 ml of the rodine solution of we multiply the remainder from the above subtraction by 0 967 we will obtain the number of mg of acetone and acetoacetic acid, expressed as acetone in the volume of urine taken for analysis

¹⁴¹ Shaffer and Marriott J Biol Chem 16 265 (1915)

[&]quot;The amount used depends upon the expected yield of \$-hydroxyl utyric acid In urines which give a strong ferric chloride reaction f c acetoacet c and or when a to 10 g or more of β-hydroxybutyric acid are expected it is unnecessary to use more than 20 to of ml of unner llowever m case only a trace of \$\tilde{\ell}\$ hydroxy butyne acid is especially volume should be much larger as indicated. Under all conditions the amount specialise is sufficient for distinct of the conditions and the conditions are sufficient for distinct of the conditions and the conditions are sufficient for distinct of the conditions are sufficient for distinct of the conditions are sufficient for distinct of the conditions are sufficient for distinct of the conditions are sufficient for distinct of the conditions are sufficient for the conditions are suffi is sufficient for duplicate determinations. It is desirable to use such a volume of urine as

contains the proper amount of β-hy droxybuty rie acid to yield 20 to .00 mg. of acetone If the urine contains I tile or no sugar only half the amount or less of lead acetate

In many instances when a high degree of accuracy is not required this reductillation n ay be om thed and the first dust llate tetrated directly. The results so obtained are slightly helper the three lates are slightly helper the three lates are slightly and the slightly helper the three lates are slightly helper the three lates are slightly helper the three lates are slightly and the slightly are the slightly a higher than those after redustulation from alkali. The object of the redustilation is to get rid of fatty acids of which tormic seid is one of the most troublesome.

denser fitted with a delivery tube drawn out to a fine tip. An all-alass apparatus is preferred, the joints being lubricated with water; if this is not available, connections may be made with well-fitting cork (not rubber) stoppers. As receiver for the distillate, use a 15-ml. graduated centrifuge tube, so arranged that the enlarged portion of the delivery tube reasts on the rim of the centrifuge tube, acting as a cover for it, and the fine tip of the delivery tube reaches just to the bottom of the receiver. Place a minimal amount of water in the receiver, to cover the tip outlet. Apply heat with a microburner to the contents of the distilling flask, slowly at first to prevent excessive hubbling in the receiver, and distif over a volume of distillate equal

to one-third or more of the original volume. Remove the receiver, rinsing off the tip of the delivery tube with a little water in the process, measure the total volume in the receiver, or dittle with water to a definite volume, and

mix by Inversion.

For color development, transfer 0.1 ml, of salicy lic aldehyde to a test tube graduated at 5 and 10 ml. Add 2 ml. of distillate, followed by 1.5 ml. of saturated potassium hydroxide solution from a huret with a fine glass tip. Nix the contents of the tube by several churning motions with a footed glass rod, leave the rod in the tube, and allow to stand 20 minutes at room temperature. Finally add either (A) water to the 10-ml. mark, or (B) squeous alcohol to the 5-ml. mark, rinsing and removing the rod in the process. Mix by tapping or inversion. Choice between procedures A and B will depend largely upon the amount of acetone present; A gives less color than B, and is for larger amounts of acetone. This choice can usually be estimated roughly by inspection during the period of color development. For many purposes one procedure can be used consistently; both are described to permit the accurate conserved of a wide ranse of acetone concentration.

filtered. Transfer an aliquot of the filtrate (the amount depending upon the 8-hydroxybutyric acid content of the urine) to a distilling flask similar to that used for the acetone determination, but in addition fitted with a dronping funnel. Make un the volume in the flask to not less than 30 mi. by adding water if necessary, and acidify with three to four drops of 1:1 sulfurio acid Distill off one-third or more of the original volume as described for the acetone determination, to get rid of preformed acetone and acetoacetic acid Discard the distillate: It cannot be used for simultaneous acetone determination since some acetone is lost by the treatment with calcium hydroride and copper sulfate. Piace a 200-ml, round-bottomed flask in position as receiver for the distillate, closed and containing sufficient water to cover the tin of the delivery tube as described under "Determination of Acetone," above. Bring the residual solution in the flask (volume of approximately 20 to 60 ml) to a holl and add through the dropping funnel 15 ml of 1:1 sulfuelc acid and 10 ml, of 0.2 per cent potassium highromate solution, by drops, during the first five minutes of distillation, followed by 25 ml, of high tomate during each of the part two five-minute periods. Regulate the rate of distillation so that 50 to 85 ml, of distillate are obtained in 15 minutes.

Measure the volume of distillate, or dilute to a known volume with water and mix, and determine the acetone content of a 2-ml, portion as described above.

CALCITATION For colorimetric measurement

 $\frac{\text{Rending of Standard}}{\text{Rending of Unknown}} \times S \times \frac{\text{Distillate Vol}}{2} \times \frac{100}{\text{Urine Vol}} = \frac{\text{mg acetone per}}{100 \text{ ml urine}}$

where S is the rectone content of the standard in mg, and the volumes of distillate, and unne whose acctone is contained in the distillate, are measured in an \mathbb{N} . The stand ard should have an acctone content approximating that of the distillate If this is not known, or will vary widely from one unknown to another, a series of five standards may be prepared which will permit the accurate estimation of any concentration of acctone in the distillate from 0.05 to 200 mg per cent (0.001 to 40 mg per 2 ml.) First prepare dilute standard acctone solutions containing 0.1, 1.0, and 5.0 mg per cent acctone, by dilution of the stock standard Set up five test tubes, each containing 0.1 ml of selective addeby de Add to each the amount of a particular standard as indicated in the table, followed by 1.5 ml of potyssium hydroude solution and subsequent treatment for color development exactly as described for the analysis of an unknown Dilute each standard after color development to the volume and with the solvent indicated in the table. The S value for each standard, to be used in the calculation, is given in the last edginn of the table.

Fanal Volume

need be prepared. If the color of the unknown in the alcohol solvent is more than twice that of Standard 3 or in the water solvent more than twice that of Standard 5, it may be diluted with the same solvent until approximate color match with a standard is obtained, and then read Results in this event must be multiplied by the dilution. Dilution up to 200-ml final volume is permissible

For photometric measurement the acetone content in the 2 ml of distillate taken for analysis is established by reference to a calibration curve prepared previously from standard acctions solutions. The calculation is then

$$\begin{array}{ll} \text{Mg acctone in} \\ \text{2 ml distillate} \times \frac{\text{Distillate Vol}}{2} \times \frac{100}{\text{Urine Vol}} = \frac{\text{mg acctone per}}{100 \text{ ml urine}} \end{array}$$

To prepare a calibration curve for procedure A (water solvent 10 ml volume). 2-ml portions of standard acctone solutions containing from 0.0 to 0.15 mg of acctone give a satisfactory curve relating density and concentration at 520 mg and 1 em solution depth (or its equivalent). For procedure B (alcohol solvent 5 ml volume) the corresponding range is 0.0 to 0.0 mg of acctone If the unknown is beyond the range of the curve repeat the analysis using a smaller aliquid of distillate (or dibited distillate) made up to 2 ml with water, and correct the calculations accordingly. In using a calibration curve for photometric measurement the experimental conditions during an analysis (time of standing before and after dibition temperature, etc.) should be uniform and should reproduce as far as possible those used when the curve was constructed. For accurate results the curve should be checked at intervals particularly if new reagents are prepared and reestablished if necessary.

Results obtained by the procedure in which β hydroxy bittyric acid is not determined (Determination of Acetone and Acetoneetic Acid) represent the preformed acetone present as well as acctone equivalent to the acetoacetic acid present which is decomposed into acetone equivalent of the acetoacetic acid present which is decomposed into acetone equivalent of the β hydroxybutyric acid present For factors used to obtain the content of β -hydroxybutyric acid present For factors used to obtain the content of β -hydroxybutyric acid itself ace previous methods

INDICAN

Methods, Indican (indoxylsulfune acid) is usually determined by oxidation and condensation to form indigo or similar substances, followed by extraction with chloroform or other suitable solvents, and colorimetric or photometric estimation in terms of a standard. Earlier methods employed a standard solution of indigo. This type of standard has been criticized by Meiklejohn and Cohen, 119 type of standard has been criticated by Meiklejohn and Cohen, 119 who state that the final color intensity obtained from urine may be deeper than that corresponding to a saturated solution of pure indigo in the same solvent. These authors describe a photometric modification of the method of Sharlit, 100 which appears to give satisfactory results and which is based upon calibration of the photometer with standard indican solutions. In Sharlit's procedure, developed for visual colorimetry standard indican solutions may also be used, but the author likewise describes an artificial standard containing cobalt sulfate, whose indican equivalence is defined. Kumon''il lass described a procedure claimed to be outle specific, based upon the

¹⁴ Merklejohn and Cohen J Lab Clin Med 27, 949 (1941 1942) See also Townsend

Dod 23 809 (1937 1938) 178 Sharlit J Biol Chem 99 537 (1932) 171 Kumon Z physiol Chem 231 205 (1935)

color reaction between indican and ninhydrin. It is not felt practical to describe any of these procedures here, because of the difficulty of obtaining pure indican for standardization purposes ¹⁷² The reader is referred to the original papers for details

Interpretation. The daily exerction of indican ranges from 10 to

20 mg per day in normal individuals

Indican is apparently formed within the body from indole produced in the intestinal lumen by bacterial action, there is no good evidence that either indole or indican are intermediates in tryptophan metabolism by animal tissues. In normal individuals, variations in indican exerction appear to be dependent mainly upon the diet, a ment diet increasing exerction while a milk or carboby drate diet decreases exerction. Pathologically the greatest increases are found in disorders involving increased putrefaction and stagnation of intestinal contents. Bacterial decomposition of body protein, as in gangrene, putrid pus formation, etc. gives rise to increases.

PHENOLS

Method of Volterra *** Principle The urine is distilled from slightly alkaline solution to obtain the free volatile phenols in the distillate After acidification, a second distillate is obtained, this represents the conjugated volatile phenols present Ether extraction of the remaining fluid separates the aromatic hydroxy acids from residual phenols. The significance of these various fractions is discussed under "Interpretation" Each fraction after proper preparation, is treated with the phosphotungs the phosphotung discussed under the standard phenol solution.

Procedure 174

(a) FREE VOLATILE PHENOLS Transfer 10 ml of the well mixed 24-hour sam ple of urine to a 250 ml distilling flask fitted with a condenser Add 150 to 175

¹⁷² For method of preparing indican from the urine of dogs fed indole see Ellinger Z physiol Chem 38 178 (1903)

¹⁷² Volterra Am J Clin Path 12 525 580 (1942)

¹⁷⁴ Reagents Required Silver Lactate Solution 3 per cent silver lactate in 3 per cent lactic acid

iactic acid

Colloidal Iron Fisher Scientific Company s Dialyzed Iron 5 Per Cent is satisfactory

deed Sedium Chloride Solution To 1 liter of a saturated solution of sodium chloride add 10 ml of concentrated by drochloric acid

Phenol Reagent (Tohn and Cuccalteu J Biol Chem 73 697 (1927)) Into a 1500-ml Irornee flask introduce 100 g of sodoum tungstate NasWO, 21 f O 25 g of sodoum molybdute Nas WO, 21 f O 25 g of sodoum molybdute Nas WO, 21 f O 25 g of sodoum molybdute Nas WO, 21 f O 25 g of lathium sulfate 50 ml of water and a few drops of bromme Boil the maxture for 15 minutes with out condenser to remove excess bromme Cool dutte to 1 liter and filter The reagent should have no greenish tint Protect from dust Didute a portion with an equal volume of water before use

Star dar! Phenol Sol ton Prepare and standardize a stock solution of phenol containing I mg of pieuol per ml as follows Desolve at latte over I g of ery stallized phenol in a liter of 0.1 N by drochloric acid Transfer 25 ml of this solution to a 250-ml flask add 50 ml of 0.1 N sodium hydroxide heat to 6.3° C add 2.5 ml of 0.1 N sodium selection temperature for 30 or 40 minutes, 1dd 5 ml of concentrated hydrochloric acid and titrate the excess of soline with 0.1 N thosulfate solution Field ml of 0.1 N sodiue solution used up correspond at 0.1 557 mg of phenol On the br is of the result dilute the remainder of the phenol solution with 0.1 N arch 150 mg of 10 ml of 0.1 N arch 150 m

ml. of water, followed by sufficient sodium hicarbonate solution to render the solution alkaline to littims. Distil, collecting the distillate until 30 to 40 ml. have been obtained. Stop the distillation and measure the volume of distillate. Use a 10-ml. portion for the color reaction, as described below.

- (b) CONIGATEO VOLATILE PIRNOIS When the contents of the distilling flash have cooled somewhat, add sufficient dilute sulfuric acid to creader distinctly acid to Congo red, and start the distillation again. Collect 100 to 120 ml. of distillate, then test for completeness of distillation by collecting a separate 5-ml. portion of distillate and treat this by the phenol color reaction described helow, using proportionately reduced amounts of reagents. If an evident color reaction is obtained in this test sample, add 50 to 100 ml. of water to the distilling flask and continue distillation until volatile phenol no longer distill over. Only in exceptional cases must more than about 150 ml. of distillate he obtained. Measure the volume of distillate, mlx, and ust a 10-ml, portion for the color reaction.
- (c) TOTAL ARGMATIC HYDROXY ACIDS Transfer the acid solution in the distilling fiask to a 250-ml, separatory funnel, using water for rinsing and diution to about 100 ml, and shake for 3 minutes with 30 to 40 ml, of petroleur ether. Draw off the aqueous layer into a second separatory funnel (discard the petroleum ether layer) and shake twice with 30-ml, portions of ordinary ether, removing and combining the ether extracts, and saving the residual aqueous fluid. Wash the combined ether extracts by shaking with water, draw off the water, transfer the ether extract to an evaporating dish, and evaporate off the ether on a steam bath. Just before evaporation is complete, add about 5 ml, of water, and then complete the removal of the ether. Transfer the aqueous fluid remaining to a graduated cylinder and make up with rinsings to 10 ml. Nix, dilute 1 ml, to 10 ml, with water, and use this diluted portlon for color development.
 - (d) BITSIDUAL PHENOLS Dilute the residue from the ether extraction to 200 ml. with water. To a 10-ml portion, add 1.0 to 1.5 ml. of silver lactate solution, and one to two drops of colioidal iron. Shake, dilute to 20 ml. with water, allow to stand 15 mlnutes, and filter. To 10 ml. of filtrate add 1.0 to 1.5 ml. of acid sodium chloride solution, dilute to 20 ml. with water, mix, and filter. Use 10 ml. of filtrate for the color reaction

Color Reaction To 10 mi. of unknown in a test tube, add 0.5 mi of diluted phenol reagent, followed by 2 ml. of 20 per cent sodium carbonate solutions this by shaking, and alter 20 to 30 seconds place the tube in a bolling water bath lor exactly I minute Remove and cool in running cold water. Compare the color with that obtained by treating 10 ml. of standard phenol solution (containing 0 03 mg. of phenol) by the same procedure, at the same time.

CALCULATIONS

Volatile Phenols (Free or Contugated)

Reading of Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard | Standard |

Results on the distillate from (a) gave the free volatile phenol content and from (b) the conjugated volatile phenols (both expressed as phenol). From the volume of the unner the total output per 24 hours can be calculated.

working standard dilute 0.3 mL of the stock standard to 100 ml with water and mis. This solution contains 0.03 mg, of phenol in 10 ml and is made up fresh at the time of

Aromatic Hydroxy Acids

```
Reading of Standard Reading of Unknown X mg phenol x 10 = mg aromatic hydroxy reads (as phenol) in ml urine used
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Multiply results by 1 5 to express in terms of p-hydroxyphenylacetic acid instead of phenol It may be necessary to dilute the 10-ml aqueous solution of ether-soluble material of which 1 ml is used for analysis to a greater volume to get color match with the standard in which case the value 10 in the formula is replaced by the volume after dilution

Residual Phenols

```
Reading of Standard Reading of Unknown X mg phenol X 80 = mg residual phenols (as phenol) in ml urine used
```

Interpretation. Results on normal and pathological urine by this method indicate that free volatile phenois (phenoi, p-cresol, etc.) are ordinarily present only in traces if at all, and these findings are in agreement with the results of others (Deichmann and Shafer, 175 Schmidt 176) using different methods. Conjugated phenois range in amount from 20 to 70 mg per day normally, and do not appear to be significantly altered in amount by diot or fasting at least in short experiments Pathologically, increases in conjugated phenois have been noted in conditions associated with extensive tissuo destruction and in intestinal obstruction. Ingestion of phenol or henzene likewise leads to increases. Aromatic hydroxy acids (p-hydroxyphenylacetic acid, p-hydroxyphenylpropionic acid, p hydroxyhenzoic acid, p-hydroxyphenyllactic acid, and possibly p-hydroxyphenyl pyruvic acid) are excreted normally in amounts ranging from 50 to 90 mg (as p-hydroxyphenylacetic acid) per day Pathological variations have not been extensively studied According to Schmidt (loc cit), about two-tlurds of the total aromatic hydroxy acid excretion is normally in the free form and one-third in the conjugated form. The significance of the "residual phenol" analysis is obscure Values ranging from 150 to 500 mg per day are found normally, but it is generally recognized that many nonphenolic substances, such as imidazoles, etc., are included in this fraction, rendering interpretation difficult Values for residual phenols as obtained by the method described here correspond roughly to the total phenol values of the admittedly nonspecific method of Folin and Denis 177

SULFONAMIDES (SULFA DRUGS)

Method of Bratton and Marshall as Principle The principle is the same as that applied to the determination of sulfonamides in blood (p. 657). The protein free urine is treated with nitrous acid to diazotize any free sulfonamide present excess nitrous acid is destroyed and the diszotized sulfonamide is coupled with N (1 naphthyl) othylenediamine to form a stable red color which is then compared with a standard treated in the same way. Total sulfonamide is determined after hydrolysis with neid The difference between free and total sulfonamide represents acetylated sulfonamide

¹⁷¹ Deichmann and Shafer Am J Clin Path 12 129 (1912)

¹⁷⁴ Schmidt J Biol Chem 145 533 (1942) 177 Folin and Denis J Biol. Chem 22 205 (1915) 178 Bratton and Marshall I Biol Chem 128, 537 (1939)

Procedure "Transfer 1 ml of urine to a 25 ml volumetric flask and dilute to the mark with water Mix, and place 2 ml of the diluted urine in a small flask From a buret add 30 ml of water, followed by 8 ml of 15 per cent tricholoroacetic acid Mix and filter

Free Sulfonamide Transfer a 10 ml portion of the trichloroacetic acid fil trate to a small flask or wide test tube, and treat with nitrite, sulfamate, etc. eractly as described for a blood filtrate (p. 658)

Total Sulfonamide Transfer a 10-ml portion of filtrate to a test tube or other container graduated at 10 ml, and add 0 5 ml of 4 N hydrochloric acid Place in a boiling water bath for 1 hour, cool, and make up to 10 ml with water Continue with treatment with nitrite, sulfamate, etc., as for free sulfonamide

Compare the final color obtained against a suitable standard as described for blood analysis, the same standards being satisfactory If the urine is unusually low or high in sulfonamide content, the determination is repeated on a more satisfactory dilution of the sample in every case, however, the final 10 mi portion analyzed should contain 3 per cent trichioroacetic acid The calculation is the same as for blood analysis, except that the dilution of the urine (I e, in the present instance the dilution is 500) replaces the value 20 in the blood calculations Either a colorimeter or a photometer may be used, as with blood

The procedure as described provides for unne containing protein as well as protein free urine If the urine is known to be free from protein the treatment with trichloroacetic acid may be omitted The urine is diluted so as to contain from 1 to 2 mg per cent of sulfonamide, and then 50 ml of this diluted urine, plus 5 ml of 4 \(\) hydrochlone acid are diluted to 100 ml with water Free sulfonamide is determined on a 10-ml portion of this final dilution as with a blood filtrate Total sulfonamide is determined on a 10-ml portion heated without further addition of acid, made up to volume, and the analysis continued as above

Interpretation. As with blood, the sulfonamide content of urine may vary between wide limits. The procedure as described is satisfactory for urines containing from 50 to 500 mg. per cent. If smaller or larger amounts than this are present other suitable dilutions must be made.

Sulfonamides found in the urine may be either in the form of the free drug its acetylated derivative, "so or oxidized forms possibly combined with glucuronic acid." Only the acetylated derivative does not respond to the colorimetric procedure, the difference between free and total sulfonamide therefore represents acetylated sulfonamide The proportion of free drug to its various derivatives depends upon a number of factors of which the nature of the sulfonamide itself is perhaps most important, sulfanilamide is relatively little acetylated as compared to sulfadiazine, for example Chincally, the sulfonamide content of the urine is of importance in at least two respects, these include the possible formation of

^{***} The reagents required are the same as those described on p 658 for the determination of sulfonamides in blood

¹⁰⁴ Marshall Bration and Litchfield Science 28, 597 (1938)

¹⁹¹ Scudi Science 91 486 (1940) also Scudi and Jelinek J Pharmacol 81 218 (1944)

urinary calculi by the insoluble and precipitated drug, and the possibility of renal damage and hematuria associated with the deposition of crystalline sulfonamide or acetylsulfonamide in the renal tubules Studies on these two possible manifestations of sulfonamide exerction have shown that the various sulfonamides and their derivatives differ significantly in their solubility in urine and in their propensity to precipitate out, either in the renal tubules or in the urine itself; in general, the therapy consists of maintaining an alkaline urine, since the compounds responsible are more soluble at alkaline reaction. Thus far, only the free drug and its acetylated product appear to be involved here; other exerctory forms of the sulfonamides, where they have been recognized, are quite soluble,

UROBILINOGEN

Method of Wallace and Diamond;182 Principle. A series of dilutions of urme is carried to the point where the red color resulting from the reaction between urobilinogen and Ehrlich's aldehyde reagent is just discernible. This method is regarded as more accurate than the spectroscopic method of Wilbur and Addis 183 A more extended, and probably more accurate, method for the determination of problingens in urine and feces is described by Watson et al 184 It has been shown that both mesobilirubinogea and stercobilirubinogen give the Ehrlich aldehyde reaction. Interfering substances can be extracted with petroleum ether

Procedure. One ml. of Ehrlich's aidehyde reagent185 is added to 10 ml. of undiluted urine and allowed to stand i to 3 minutes. An idea as to the quantity of urobilingen is gained by noting the rapidity and intensity of color development. Dilutions are not carried out if the color remains a light red (normal values). For higher concentrations, prepare a series of dilutions of the urine from 1:10 to 1:200, or higher, as indicated by the preliminary test. Tan water may be used, but should not be too cold. Add 1 ml. of the reagent to each dilution and after 3 to 5 minutes note the highest dilution that shows a faint pink discoloration. Express the result in terms of this dilution. The test is best performed in daylight but in the absence of bright sunlight.

Interpretation. The appearance of the color in dilutions up to 1:20 may be regarded as normal. Constipation may produce a temporary rise in urobilinogen. This substance is believed to originate by reduction of the bile pigment. The latter may be extrahepatic as well as hepatic in origin (see p. 592). Excessive amounts of urobilinogen are, therefore, excreted in diseases of the liver and biliary tract, including toxemias of pregnancy, infectious diseases, and alcoholic intoxication, and in hemolytic diseases, including poisoning by lead, sulfonal, mushrooms, and hemolytic poisons in general 158 It is claimed that urinary probilingen affords no accurate index to blood decomposition; according to Miller.

Wallace and Diamond Arch. Internal Med., 35, 698 (1925).
 Wilbur and Addis Arch Internal Med., 13, 235 (1914).

¹⁴⁴ Watson, Schwartz, Sborov, and Bertie: Am. J. Clin. Path., 14, 605 (1944); Watson and Hawkinson, ibid , 17, 108 (1947).

¹¹⁸ Sulfonamides in the urine will interfere since they likewise react with Ehrlich's aldehyde reasont. If they are present, acidify the urme and shake with petroleum ether. Shake the petroleum ether extract with aqueous alkah and carry out the test on the final neutralized aqueous extract.

Singer, and Dameshek, 187 however, the fecal output of urobilinogen is of value in this connection

OXALIC ACID

Earlier methods for the determination of oxalic acid in urine were based on the precipitation of calcium oxalate from a large volume of urine Difficulties in obtaining satisfactory checks were attributed to the effect of magnesium, phosphate, and sulfate ions upon the solubility and rate of crystallization of calcium oxalate. The following procedure is directed toward correcting the deficiencies in the older methods

Total Oxalic Acid (Method of Pouers and Lesatiniss) Principle The unne is acil hydrolysed (to decompose any oxalium acid) and the oxalic acid separated from interfering ions by extraction with ether. The extract as a acid cum oxalate and dissolved in acid permanganate solution and back titrated with thosulfate.

Procedure An all glass continuous liquid liquid extraction apparatus is used, as shown in Fig 256 This consists of a 25 ml test tube in which is

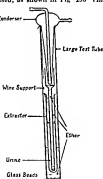


FIG 256 ALL-GLASS CONTINUOUS I IQUID-I IQUID EXTRACTION APPA-

Courtesy Powers and Levat n.

placed a funnel shaped tube made from half a 25 ml pipet having an over all length of about 200 mm The stem end is sealed and 4 holes punched through the sealed end with hot wire The tube is sus pended to an outer tube (25 × 300 mm) by means of a nichrome wire around its lip or through a small hole near the top of the inner tube This also serves to separate the cold-finger condenser slightly thus providing a vent

Of the well mixed 24-hr urine sample (preserved if necessary with a few ml of formaldehyde), transfer 25 ml to a volumetric flash, add i ml concentrated hydrochloric acid, mix by inversion and heat by immersion in boiling water for 30

minutes Cool and filter

Into the outside tube of the extraction apparatus place several glass beads, 2 ml of water and 25 ml of ether Introduce 10 ml of the acid hydrolyzed urine fil trate into the Inside tube and assemble the apparatus as shown in Fig. 256 Extraction may be carried out on an electric hot plate covered with an asbestos sheet in which a 15 cm hole is cut, the walls of the extractor being thus protected from overheating and loss of ether Dur-

ing the extraction a good flow of cold water must be passed through the condenser and the heating regulated so that the ether drips at the rate of about

iii Miller Singer and Dameshek Arch Internal Med 70 722 (1942)
Powers and Levatin J Biol Chem. 154 207 (1944)

100 drops per minute, if necessary, add a little more ether. (Warning—avoid cetting ether vapors near hot plate!) Extract for 6 hours.

Disconnect the apparatus. Lift the Inside tube, rinsing it down with 2 mi. of ethanoi from a pipet before removing. Add 1 mi. of 2 per cent acetic acid to the contents of the outside tube, shake by twirling to mix the water and ether layers (the presence of water is necessary to avoid oxalic acid loss on evaporation to dryness) and drive off the ether by Immersion in a water bath at 70° C. while continuously shaking.

Transfer the aqueous residue into a 15-ml. centrifuge tube by means of an aspirating device (see Fig. 183, p. 695). The tip of the centrifuge tube should be drawn out slightly to an inside diameter of 1 mm. so as to facilitate the handling of the small precipitate. Rinse down the wails of the outside tube with two 2-ml. portions of ethanol, transferring each rinsing to the centrifuge tube.

Add 5 mi, of 10 per cent caicium chloride to the contents of the centrifuge tube and stir with a gentle current of air introduced through a fine capillary tube. Overlay with 2 ml, of acid-ethanol solution (60 ml, ethanol, 10 ml, 2 per cent acetic acid, 20 ml, water) which prevents surface accumulation of the oxalate precipitate. Let stand overnight. Centrifuge at 2,000 r.p.m. for 30 minutes, decant, and invert on filter paper to drain. Rinse down the wall of the centrifuge tube with 2 ml. of acid-ethanol solution and thoroughly break up the precipitate with a fine glass rod. Remove the rod after rinsing it with 3 ml. of acid-ethanol solution. Centrifuge, decant, and drain as above. Drive off residual alcohol by heating in an oven or bolling water bath for a few minutes.

Add 1 ml. of 20 per cent sulfurle acid and 0.5 ml. of 1 per cent manganess sulfate. Break up the precipitate with a fine glass rod which is rinsed with a few drops of water before removal. Add exactly 3 mi. 0.01 N potassium permanganate. Stir with air current and let stand 8 10 minutes. Add 0.5 ml. of 10 per cent potassium iodide, mix by twirling, and follow with 4 drops of soiuble starch solution (see Appendix) and 2 drops of a saturated solution of barium hydroxide. (The barium sulfate precipitate facilitates visualization of the end point of the titration.) Titrate the excess permanganate with 0.01 N sodium thiosulfate from a 5 ml. microburet. Read to nearest 0.01 ml.

A titration reagent blank should be run in parallel. For this purpose distilled water may be used in place of the oralate solution.

CALCULATION.

In the following equation t is the mi of 0 01 N thiosulfate used in the back titration, b is the mi used in the reagent blank.

$$0.45(3-t-b) \times \frac{26}{25} \times \frac{24$$
-hour volume = mg total oxalic acidino in 24-hour urine

Interpretation. From 15 to 50 mg. of oxalic acid is exercted by a normal adult on an ordinary mixed duct. It arises from oxalates of the food ingested and from fat and protein metabolism. It is increased by the ingestion of apples, grapes, cabbage, etc., although most of the ingested oxalate is destroyed. It is increased in disturbances of metabolism associated with decreased oxidation, according to certain observers. The term "oxaluria" has been largely a misnomer.

SULFUR190

GRAVIMETRIC PPOCEDURES

I Total Sulfates (Folm's Method) Principle The sulfure acid of the conjugated sulfates uset free by boiling with acid. The total sulfates are then precipitated with barum chloride.

Procedure. Place 25 ml of urine in a 200 to 250 ml Erlenmeyer flask, add 20 ml of dilute hydrochloric acid*** (I volume of concentrated HCI to 4 volumes of water), and gently boil the mixture for 20 to 30 minutes. To minimize the loss of water by evaporation, the mouth of the flask abould be covered with a small water lighas during the boiling process. Cool the flask of 72 to 3 minutes in running water, and dilute the contents to about 150 ml by means of cold water. Add 10 ml of a 5 per cent solution of barium chloride allowly, drop by drop, to the cold solution **The contents of the flask should not he stirred or shaken during the addition of the barium chloride. Allow the mixture to stand at least 1 hour, then shake up the solution and filter it through a weighed Gooch crucible **

Wash the precipitate of BaSO, with about 250 ml of cold water, dry it in an air bath or over a very low flame, then ignite," cool, and weigh

CALCULATION Subtract the weight of the Gooch crucible from the weight of the crucible and the BaSO precipitate to obtain the weight of the precipitate. The weight of S in the volume of urine taken may be determined by means of the following proportion.

Viol wt Baco, g Baso, = Viol wt S z(g S)

Representing the weight of the BaSO, precipitate by y and substituting the proper molecular weights we have the following proportion

233 43 y = 3206 x(g S in the quantity of urine used)

Calculate the quantity of S in the 24 hour specimen of unne

Interpretation. The total sulfate excretion (ethereal and morganic sulfates) by a normal adult on a mixed diet is usually between 0.6 and

- 2 g. S with an average of about 1.0 g The sulfuric acid is derived but to a slight extent ordinarily from ingested sulfates, being mainly dependent on the sulfur of the protein metabolized, and will consequently vary widely with the level of protein metabolism. From 75 to 95 per cent of the total sulfur of the urine is ordinarily found as sulfate, the proportion being greatest on a high-protein diet. The sulfate excretion is increased in all conditions associated with increased decomposition of body protein as in acute fevers and decreased whenever there is a decrease in metabolic activity.
- 2. Inorgonic Sulfotes (Folin's Method): Place 25 ml. of urine and 100 ml. of water in a 200- to 250-ml. Erlenmeyer flask and acidify the diluted urine with 10 ml. of dilute hydrochloric acid (1 volume of concentrated HCl to 4 volumes of water). In case the urine is dilute, 50 ml. may be used instead of 25 ml. and the volume of water reduced proportionately. Add 10 ml. of 5 per cent barlum chloride slowly, drop by drop, to the cold solution, and from this point proceed as indicated in the method under "Total Sulfates," above.

Calculate the quantity of inorganic sulfates, expressed as S, in the 24-hour urine specimen.

CALCULATION. Follow the directions given under Total Sulfates, above

Interpretation. On an average, about 90 per cent of the total sulfates of the urine exists as inorganic sulfates, but the proportion of the sulfates existing in this form varies widely, being greater on a high protein diet than on a very low protein diet. The amount varies with the total sulfates (see above).

Ethereol Sulfotes (Folin's Method): Principle. The inorganic sulfates are removed with barium chloride and the coopingated sulfates then determined after hydrolysis.

Procedure. Place 125 ml. of urine in an Erlenmeyer flask of suitable size, dilute it with 75 ml. of water, and acidify the mixture with 30 ml. of dilute hydrochloric acid (1 volume of concentrated HCl to 4 volumes of water). To the cold solution add 20 ml. of a 5 per cent solution of barium chloride, drop by drop. 1th Allow the mixture to stand about one hour, then filter it through a dry filter paper. Collect 125 ml. of the filtrate and boil it gently for at least one-half hour. Cool the solution, filter off the precipitate of BaSO₄, wash, dry and ignite it according to the directions given on p. 946.

CALCULATION The weight of the BaSO, precipitate should be multiplied by 2 since only one-half (125 ml) of the total volume (250 ml) of fluid was precipitated by the burnum chloride. The remaining calculation should be made according to directions given under "Total Sulfates," above

Calculate the quantity of etheresi sulfates, expressed as S, in the 2t-hour urine specimen.

¹¹ See footnote 192, p 946

¹⁰ This precipitate consists of the morganic sulfates. If it is desired, this has 0, precipitate may be collected in a Goods cruable or on an ordinary quantitative filter paper and a determination of morganic sulfates made, using the same technique as that suggested above. In this way we are enabled to determine the morganic and ethereal sulfates in the same sample of urne.

Interpretation, The exerction of ethercal sulfates (expressed as S) varies ordinarily from 0.04 to 0.1 g per day comprising from 5 to 15 per cent of the total sulfur excretion. The absolute amount of ethercal sulfate increases with increase in the protein of the diet and particularly with increase of putrefactive processes in the intestine or elsewhere The amount excreted cannot, however, be taken as an index of the extent of intestinal putrefaction

4 Total Sulfur (Benedict's Methodis) Principle The urine is evaporated and graited with a solution of copper pitrate and potassium chlorate. Organic matter is thus destroyed and all unoxidized sulfur is oxidized to the sulfate form and can be readily precipitated with barium chloride in the usual manner. The method is very convenient and accurate

Procedure Ten ml of urine is measured into a small (7 to 8 cm) porceiain evaporating dish and 5 ml 199 of Benedict's sulfur readent 100 is added The contents of the dish are evaporated over a free flame which is regulated to keep the solution just below the boiling point, so that there can be no loss through spattering When dryness is reached the flame is raised slightly until the entire residue has blackened. The flame is then turned up in two stages to the full heat of the bunsen burner and the contents of the dish thus heated to redness for 10 minutes after the black residue (which first fuses) has become dry This heating is to decompose the last traces of nitrate (and chiorate) The flame is then removed and the dish allowed to cool more or less completely Ten to 20 mi of dilute (1 4) hydrochloric acid are then added to the residue in the dish which is then warmed gently until the con tents have completely dissolved and a perfectly clear, sparkling solution is obtained This dissolving of the residue requires scarcely 2 minutes With the aid of a stirring rod the solution is washed into a small Erlenmeyer flash, "1 diluted with cold, distilled water to 100 to 150 ml 10 ml of 10 per cent barium chloride solution added drop by drop and the solution allowed to stand for about an hour It is then shaken up and filtered as usual through a weighed Gooch crucible Controls should be run on the oxidizing mixture

CALCULATION Follow directions given under Total Sulfates p 946 Calculate il e quantity of sulfur expressed as S in the 24-hour urine specimen

Interpretation The total sulfur excretion averages about 10 g pcr day, expressed as S It runs more or less parallel with the decomposition

For determinat on of total sulfur in other biological material, see the method of Stock h lm and Koch (p 643) See also Pollock and Partansky Ind Eng Chem Anal Ed 6 330 (1931) 11 Benedict J Biol Chem 6 363 (1909)

in If the urine is concentrated the quantity should be slightly increased see Crystallized copper mirate sulfur free or of known sulfur cor tent 200 € Sod um or pr tawium chlorate 50 g

Distilled water to 1000 ml Denis suggested the use of the following sclution Copper n trate

²⁵ g best um ett e fe 25 g Amn nim r trate to e

of endogenous and exogenous protein and a definite ratio between the excretion of total N and total S might be expected, however, no constant value can be given See "Total Sulfates, p 946"

5 Total Sulfur (Osborne-Folin Method²²³) Principle This method depends on the destruction of organic matter by means of sodnim peroxide It is employed particularly for the determination of sulfur in foods and feese Benedict's procedure (see above) is guider and fully as satisfactor, for urine

VOLUMETRIC PROCEDURES

6 Volumetric Determination of Ethereal and Inorganic Sulfates and Total Sulfar (Fiske's Modification of the Method of Rosenheim and Drumniond)
Principle The sulfates of the urine are precipitated by means of benadine solution, the precipitate of benadine sulfate being filtered off and the sulfure acid of the compound titrated with 0.02 N NaOH, using phenolsulforephthalein (phenol red) as an indicator This is possible because the benadine is a very weak base and its sulfate readily dissociates it is necessary that excess of HCl be avoided in the precipitation process For critical studies of this procedure see Oven²²³ and McAittrek and Schmidt ²⁴⁴ Ishin and Lieboff ²⁴⁵ have described a colorimetric method for sulfate based on diarotization of the henzidine sulfate and coupling with phenol in alkaline solution.

Procedure

(a) REMOVAL OF PROSPRATE This step is necessary for the highest accuracy, especially where the proportion of phosphorus to sulfur may be high, as in half hourly specimens of urine For 24 hour specimens where the highest accuracy is not desired it may be omitted.

Transfer to a 100 ml volumetric flask enough urine (usually 10 to 20 ml) to contain about 10 to 20 mg of sulfur as sulfate Dilute to about 50 ml with water Add one drop of phenolphthalein solution and then concentrated ammonium hydroxide drop by drop to a faint pink color Add 10 ml of 5 per cent ammonium chloride and about 15 g of finely powdered magnesium carbonate Make to matk, stopper, mix thoroughly by shaking for 1 minute, and let stand for 30 minutes Using a dry filter, filter into a dry flask. This filtrate is used for the three following determinations

(b) Inomonato Sulfatte Pipet 5 ml of the filtrate into a large pyrex test tube with flaring lip Add 2 drops of a 0 04 per cent alcoholic solution of brome-phenol blue and 5 ml of water Add approximately N IICl until the last trace of blue disappears and the solution is yellow Add 2 ml of benzidine solution 10 test stand for 2 minutes Add 4 ml of 25 per cent acctone and let stand for 10 minutes Prepare a thin mat of paper pulp in a filtration tube (see Fig 257) 207 This mat should first be washed with water and then sucked dry Filter off benzidine sulfate with very genite suction Wash down the

sides of the test tube with 1 ml. of 95 per cent acetone, transferring to filtration tube. Wash twice more with 1 ml. and finally with 5 ml. of acetone. Add about 2 ml. of water and poke the mat with a wire out through the bottom of the tube into the pyrex test tube, rinsing the wire with a few drops of water. Add 2 to 4 drops of 0.5 per cent water solution of phenol red. Titrate with 0.2 N NaOII (prepared from 0.1 N NaOII by dilution) the solution being kept



FIG 257 FILTRA-TION TUBE FOR SULPUR DETERMI-NATION (FISKE) 144

red from 0.1 N NaOll by dilution) the solution being kept hot. At the beginning of the titration the filter tube is kept suspended in the mouth of the test tube and the alkali run through the filter tube to dissolve adherent sulfate. Rinse the inner tube with 2 to 3 ml. of water, heat the solution to boiling so that the tube is further washed with condensed steam, and finally rinse the inner tube with a few ml. more of water and remove. Titrate to a definite pink color that remains after boiling.

CALCULATION The sulfate titrates like free sulfuric acid. One ml. 0.02 N NaOH is equivalent to 0.32 mg. S. Multiply the buret reading by 0.32 to get mg. inorganic S in 5 ml. filtrate.

- (c) TOTAL SLIFATE (LORGANIC AND ETHEREAL) TO 5 ml. of filtrate in a 100-ml. beaker add 1 ml of approximately 3 N liCl. lieat on a water bath to dryness and for 10 minutes longer. Transfer to a pyrex tube with lip using five 2-ml. portions of water, add 2 ml. of benzidine solution, and proceed as in the method for inorganic sulfate The calculation is the same
- (d) PTHEREAL SULFATE Subtract inorganic from total sulfate. The difference is ethereal sulfate.
- (e) TOTAL SULFUR Transfer approximately 0.25 ml. of Benedict's sulfur reagent (see p. 948) and 5 ml. of filtrate to a small evaporating dish (6 cm.). Expaprate carefully to dryness on wire gause or hot plate. Gradually increase heat, finally igniting at red heat for 2 minutes over free fiame. Let cool for 5 minutes. Add 1 ml. of 3 N liCl and evaporate to dryness at low heat. The mixture will turn from green to brown Transfer to a pyrex tube with a lip with the aid of five 2-ml. portions of water. Add 1 drop of N liCl and 2 ml. of henzidine solution. Complete the determination as in the methods above but in place of the first 1-ml. portion of acctone, in washing use 2 ml. of 50 per cent acctone The calculation is the same as above.

(f) NEUTRAL SULFUR. Subtract from the total sulfur the total sulfate as determined above. The difference is neutral or unoxidized sulfur.

Interpretation. The neutral sulfur of the urine is made up of cystine and related substances, thiocyanate, oxproteic acids, etc. It makes up ordinarily from 5 to 25 per cent of the total sulfur of the urine, or on the average 0.98 to 0.16 g. per day calculated as S. The absolute amount is fairly constant for a given individual through wide variations of protein intake, indicating that its origin is mainly endogenous, that is, that it arises principally from the decomposition of tissue protein. On this account the percentage of the total sulfur excretion existing in the neutral form may rise to 25 per cent on a very low protein diet and decrease to 5 per cent on a high-protein diet, the absolute amount remaining nearly constant. In fasting percentages as high as 70 have been noted Im many disorders, as title reuliess inner e systimuma, etc., the amount may be relatively and m some cases absolutely increased, but no fixed relations have been determined for the various conditions.

PHOSPHORUS

1. Determination of Inorganic Phosphate (Method of Fiske and Subba-Row203): Principle. Phosphate reacts with moiybdic acid to form phosphomolybdic acid On treatment with 1,2,4-aminonaphtholsulfonic acid, phosphomolybdic acid is selectively reduced to produce a deep blue color ("molybdenum blue"), which is probably o mixture of lower oxides of molybdenum. This color is then compared in a colorimeter or photometer with that obtained from a suitable standard phosphate solution treated in the same way.

Procedure, 211 Measure into a 100-ml, volumetric flash enough urine 212 to contain between 0.2 and 0 8 mg. of inorganic phosphorus (usually 1 or 2 ml.). Add water to bring the total volume to about 70 ml, followed by 10 ml. of Molybdate I reagent, Mlx by gently shaking and odd 4 ml of aminonaphtholsulfonic acid reasent. Again mix, dilute to the mark with woter, mix several times by inversion, and allow to stand 5 minutes

At the same time, transfer to a similar flask 5 ml of standard phosphate solution, containing 0.4 mg, of phosphorus, 65 ml, of water, and the same reagents that were added to the urine sample, mixing as above. Dllute to the mark with water, mix well by inversion, and allow to stand 5 minutes. For photometric measurement, prepare a blank solution by treating 70 ml. of water in 0 100-ml. flash with the same reagents that are used for the standard and unknown, dilute to the mark with water, and mix by inversion.

For colorimetric measurement, match the standard against itself at 20 mm. and compore the unknown ogainst the standard For photometric measurement, determine the density of the stondard and of the unknown in a photometer at 660 to 720 my, setting the photometer to zero density with the blank.

CALCULATION For colorimetric measurement

Reading of Standard Reading of Unknown × 0.4 = mg inorganic phosphate (as P) in the volume of urine used

An unknown reading between 5 and 40 mm is acceptable with the standard at 20 mm. if the unknown reads outside this ronge, repeat the analysis on a smaller or larger volume of urine Results are usually expressed in terms of grams (or mg) of inorganic P per 24-hour volume of urine

For photometric measurement

Density of Unknowo Density of Standard × 04 = mg inorganic phosphate (as P) in the volume of prince used in the volume of urine used

201 Fiske and SubbaRow J Biel Chem 66, 375 (1925)

200 Various other reducing agents have been proposed (see discussion on p. 630), of which stannous chloride has perhaps found most favor A method using stannous chloride has been suggested 110 Stannous chloride has the advantage of being more stable than aminonaplitholsulfonic acid solution, and of giving a more intense color, thus permitting the use of smaller volumes of urine This latter appears to be of little value in urine analysis and a disadyantage of stannous chloride is that the color produced changes in intensity on standing and may not obey Beer s law, thus requiring careful technical control, particularly in photometric measurement

210 Youngburg and Youngburg J Lab Clin Med. 16, 158 (1930)

In The reagents required and the standard phosphate solution, are described on p 631

in connection with the determination of blood morganic phosphate

212 If albumn 15 present and the addition of moly belate produces a turbidity, treat the urine with 4 volumes of 10 per cent trichloroscetic acid, stopper, shake and filter, and repeat the determination using 2 to 10 ml of filtrate If very low in phosphate so that more than 19 ml of filtrate must be used follow the method for blood filtrates (see p. 631)

In a 1-cm cuvette, and at 660 mµ the density of the standard is approximately 0 500 (see Fig 166 p 632). Under these conditions up to 0 8 mg of phosphorus may be accurately determined. If larger amounts are present or if a deeper cuvette is used the determination is carried out on a smaller portion of urine (and standard, if necessary) and the calculations are corrected accordingly.

Interpretation. The exerction of phosphoric acid is extremely variable but on the average the total output for the 24 hours is about 11 g expressed as P, mainly in the form of phosphates. The greater part of the phosphate exerction arises from the ingested food, either from the preformed phosphates or more especially from the organic combinations as phospho- and nucleoproteins. The exerction is consequently very largely dependent upon the phosphorus content of the diet. Some of the phosphorus and results from the hreakdown of the tissues of the hody, and this endogenous phosphoric acid exerction is increased in conditions of increased metabolism as in fevers. The findings in pathological conditions have been somewhat contradictory due to lack of control of diet. The so-called "phosphaturas" nearly always represent decreased acidity and not increased phosphate content of the urine. Such conditions are how ever, significant as indicating a possible tendency to the formation of phosphatic calcul

2 Turni Phosphites (Uranium Acetate Method) Principle Standard uranium acetate is rui into a measured quantity of urine until all of the phosphate has been precipitated as insoluble uranium phosphate has received uranium in indicated by a reddish coloration with potassium ferrocyanide. This method is accurate and gives practically the total phosphorus of urine maximuch as the latter exists generally almost entirely as phosphates.

Procedure To 50 ml of urine in a small beaker or Erlenmeyer flask add 5 ml of a special sodium acetate solution¹¹ and beat the mixture to the boiling point From a buret, run into the hot mixture, drop by drop, a standard solution of uranium acetate¹¹ until a precipitate ceases to form and a drop of the mixture when removed by means of a glass rod and brought into con act with a drop of a solution of potassium ferrocyanide¹¹ on a porcelain test tablet produces instantaneously a brownish red coloration. Take the buret reading and calculate the P content of the urine under examination

CALCULATION Multiply the number of ml. of uranium acetate solution used by 0 002 to determine the number of grams of P in the 50 ml of urine used. To express the

^{2.1} The sodium acetate solution is prepared by dissolving 100 g of sodium acetate in 500 ml of distilled water adding 100 ml of 30 per cent acetic acid to the solution and making the volume of the mixture up to 1 liter with water.

¹¹ Uranium Accide: Solution Desolve about \$5.0 g. of uranium acetate in 1 liter of water with the aid of heat and \$1.0 st ml. of placal aceta cancil Let stand a few days and filter Standardize against a phosphate solution containing 0.002 g of P per ml For the ruppose dissolve 13.40 g of pure air-dry sodium ammonium phosphate (As MILHFO)+411(4) in water to make a liter To 20 ml of this phosphate solution in a 200-ml better the first solution (see above) and the solution of the solution of the dism accetate solution (see above) and till the text of the transium solution to the contract of the dism accetate solution (see above) and till the except 20 ml of uranium solution are required t ml of the solution is equivalent to 0.007 g of F H stronger than this dilute accordingly and check again by tirtation.

[&]quot;A 10 per cent solution of potassium ferrory and its assistatory. Cochimeal in 30 per cent alord of may be used as an indicator it is added directly to the urine after the unstandance accetate tristion produces no further precipitate. A green color is the end creation. The use of cochimeal is more environmental but rather less accurate than the procedure involving the use of the ferrory and the second control of the contro

result in percentage of P, multiply the value just obtained by 2, e.g., if 50 ml of urine contained 0 074 g. P, it would be equivalent to 0 148 per cent

3. Total Phosphorus: Principle. Although urme phosphorus exists almost entirely as morganic phosphate and the determination as such is usually sufficient, a strictly total phosphorus determination requires the destruction of organic matter. This is brought about by heating with sulfuric acid and hydrogen peroxide. The determination is otherwise the same as for the colorimetric determination of inorganic phosphate.

Procedure. Place sufficient urine to contain hetween 0.2 and 0.8 mg. of total phosphorus (usually 1 to 2 ml.) in a large pyrex test tube (200 hy 25 mm.), add 10 ml, of 5 N sulfuric acid (or 5 ml, of 10 N acid), and a quartz chip to prevent humping. Heat over a microhurner or on an electric hot plate until the water has been driven off and a dark brown fluid remains Add 30 per cent hydrogen peroxide216 hy drops as necessary to complete oxidation of organic matter and leave a colorless solution, heating to boiling momentarily hetween drops. The hydrogen peroxide should he allowed to drop directly into the tube contents and not run down the sides of the tube. When oxidation is complete, cool slightly, add 2 to 3 ml of water, and holl for two to three minutes to ensure hydrolysis of meta- or pyrophosphate. Cool, and transfer quantitatively with rinsings to a 100-ml volumetric flash. Make up to about 70 ml, with water and add 10 ml, of a 25 per cent solution af ammonium molyhdate in water. Mix and add 4 ml of the aminonaphtholsulfonic acid reasent used far the determination of inorganic phosphate (p. 951). Dilute to 100 ml, with water and mix. The remaining procedure (including standard and calculations) is exactly as described for the determination of inorganic phosphate.

4. Total Phosphorus (Neumann's Procedure): Principle. The organic matter is destroyed by digestion with a mixture of sulfure and mitra acids or some other axidizing agent. The phosphorus is then precipitated as the phosphomolybdate and determined gravimetrically or volumetrically.

Preparation of the Solution. Pipet 10 ml. of urine (or an amount of substance containing about 10 mg. of P) into a X-peldahi Bask. Add 16 ml. of a mixture of equal parts of concentrated H-SO, and concentrated HNO,. Digest over a low flame until red fumes cease to come off. If the mixture darkens due to the charring action of the sulfuric acid, add nitric acid from a separatory funnel a few drops at a time and continue the digestion. When the mixture remains clear on evaporation to the point where white sulfuric fumes come off, the digestion is completed by heating for 10 to 15 minutes longer. Cool and transfer the solution to a 400-ml. Erlenmeyer flask with the ald of enough water to make a total volume of about 75 ml. 32.

Instead of oxidizing the material as described above, it may be ignited with magnesia to destroy organic matter. About 2 g. of the solid substance or 25 mil. of urine (previously evaporated nearly to dry ness) are mixed with a little more than an equal buil, of magnesium oxide in a porceiain dish of about 30-mi, capacity. Five mil of magnesium nitrate solution (see Appendix) are added and the mixture heated very gently at first, then gradually to bright

²³⁴ Merck s Blue Label or "Special" Reagent is satisfactory

if In the case of urine it is possible to neutralize this seid solution with ammonis make it acid with sectic acid, and titrate with uranium acctate as in the preceding method

redness. The mixture is cooled and transferred with water to a 250-ml. flask. An excess (20 to 30 ml.) of HCl are added and the mixture boiled a few minutes. Remove from the flame and add at once enough barium chloride solution to precipitate any sulfate present. Cool, make to mark, filter, and take an aliquot for analysis.

Precipitation of the Phosphomolybdate. Neutralize the solution with ammonia, make slightly acid with nitric acid, and add 15 g. of ammonium nitrate in substance (or 25 ml. of a 60 per cent solution). Heat on a water bath to 60° to 65° C. (not higher) and add 30 to 40 ml. of molybdate solution, "is str and let stand for about 15 minutes at 60° to 65°. Filter at once through a small paper, "" washing the precipitate twice by decantation with 1 per cent potassium nitrate solution, using about 25 ml. each time, attering up the precipitate well in each case, and allowing to settle. Transfer the precipitate to the filter and wash with 1 per cent potassium nitrate solution until two fillings of the filter (collected separately) do not greatly diminish the color produced with phenolphthalein by 1 drop of the standard alkall.

Titration of the Phosphomolybdate. Transfer the precipitate and filter back to the original beaker and dissolve in a small excess of 0.2 N NaOII (about 2 to 3 ml. more than required to completely dissolve the yellow precipitate). Add about 100 ml. of boiled and cooled water and a few drops of phenolphthalein as an indicator (a red color should be observed indicating excess of NaOII) and titrate the excess of NaOII with 0 1 N acid.

CALCUATION, Divide the ml of 0 I N acid used by 2 and subtract from the ml of 0 2 N NaOH used. This gives the ml of 0 2 N NaOH required. Yightipp to 0270 (the equivalent of 1 ml 0 2 N NaOH in P) to obtain the number of mg P in 10 ml of the urine analyzed. Calculate the daily output of P in this case from the 24-hour volume.

Interpretation. Nearly all of the phosphorus of the urine exists as inorganic phosphates. Consequently the total phosphorus of urine varies in the same way as total phosphates. A small portion of the phosphorus of the urine may exist in organic combination, though not in a reduced form. This organically bound phosphate may amount to from 1 to 4 per cent of the total phosphorus excretion. Lattle 18 known with regard to the compounds in which it occurs. Some glycerophosphoric acid may occur either free or as lectifus.

of ammonium hydroxide solution (sp. gr. 0 90) Let stand for some time (2 hours is usually enough), then filter and wash the precipitate with 25 per cent ammonia until practically free from chlorides Ignite to whiteness or to n gray 18h-white ash and weigh Multiply the weight of magnesium pyrophosobate thus obtained by 0.279 to get the weight of P.

CALCULATION Calculate as explained above

CHLORIDES

1 Volhard-Arnold Method Principle The urine is acidified with intric acid and the chlorides precipitated with a measured excess of standard silver intrate solution. The silver chloride formed is filtered off and in the filtrate the excess silver intrate is titrated back with standard ammonium thiocyanate solution. Ferric ammonium sulfate is used as an indicator. A red color due to the formation of ferric thio cyanate indicates that an excess of thiocyanate is present and that the end point has been reached.

Procedure, Place 10 ml of uriae in a 100-ml volumetric flash, add 20 to 30 drops of nitric acid (sp gr 1 2) and 2 ml of a cold saturated solution of ferric aium If necessary, at this point a few drops of 8 per cent solution of potassium permanganate may be added to dissipate the red color. Now slowig run in a known volume of the standard silver intrater solution (20 ml is ordinarily used) in order to precipitate the chlorine and insure the presence of an excess of silver nitrate. The mixture should be continually shaken during the addition of the standard solution. Allow the flask to stand 10 miautes, then fill it to the 100-ml graduation with distilled water and thoroughly mix the conteats. Now filter the mixture through a dry filter paper, collect 50 ml of the filtrate, and titrate it with standardized ammonium thiocyanate solution. The first permanent tinge of red-hrown indicates the end point. Take the buret reading and compute the weight of sodium chioride in the 10 ml of uriae used.

CALCULATION The number of ml of ammonium thiocyanate solution used indicates the excess of standard silver nitrate solution in the 50 ml of filtrate trimted Multiply this reading by 2, inasmuch as only one-half of the filtrate was employed, and subtract this product from the ml silver nitrate (20 ml) originally used, in order to obtain the ml of silver nitrate utilized in the precipitation of the chlondes in the 10 ml of unne employed.

To obtain the weight in g sodium ablonds in the 10 ml of urne used, multiply the ml of the standard silver intrate solution actually utilized in the precipitation, by 0 010 If it is desired to express the result in percentage of sodium chloride, now ethe decimal point one place to the right, for results in terms of grams of sodium chloride ner liter of urne, move the decimal point two places to the nebt

³ Standard silver nitrate solution may be prepared by dissolving 29 061 g of silver nitrate in 1 liter of distilled water Each mi of this solution is equivalent to 0 010 g of solume hiboride or to 0 000 g of chlorine

^{**}This solution is made of such a strength that I ml of it is equal to I ml of the standard silver nutrate solution used To prepare the solution dissolve 13 g of ammonium thocyanate NHSCN in a little less than a liter of water. In a small flask place 20 ml, of the standard silver nutrate solution 5 ml of the ferrica dism solution and 4 ml of nutrica acid (sp. gr. 12) add water to make the total volume 100 ml and thoroughly must be contents of the flask. Now run in the ammonium thiocyanate solution from a hiret until a perimanent red brown tinge is produced Thes is the end reaction and indicates that the last trace of silver nutrate has been precipitated. Take the burset reading and calculate the amount of water necessary to use in disturing the ammonium thocyanate in order that 10 ml of this solution may be exactly equal to 10 ml of the silver nutrate solution. Vake this dilution may he exactly equal to 10 ml of the silver nutrate solution. Vake

In a similar manner the weight or percentage of chlorine may be computed using

the factor 0 006 instead of 0 010 To express results in terms of milhequivalents of chloride per liter of urine multiply the ml standard silver nitrate required for the chloride in 10 ml of urine by the factor

17 1 Calculate the quantity of sodium chloride and chlorine in the 24-hour urine specimen

Interpretation. From 10 to 15 g of chlorine, expressed as sodium chloride (170 to 250 milliequivalents of chloride), are excreted per day, on the average, by normal adults. The amount is, however, closely dependent upon the chloride content of the food ingested. In fasting, the chloride excretion falls rapidly to a very minimal quantity. On high water ingestion it is increased. In pneumonia and certain other acute infectious diseases the excretion of chlorides may be markedly diminished particularly during the periods in which exudates are forming. In convolucional and with resolution of the exudates the chlorine excretion rises again. A decrease has also been noted in nephritis associated with edema

- 2 Volhard Harsey Method. Principle This procedure!³³ differs from the \0 of hard Arnold method in that the excess of silver intrate is titrated directly without filtering and tence in the presence of the alver chloride. The procedure is thus more rapid but the end point is more difficult to determine, and the results are not so exact it is therefore not recommended.
- 3 Method of Sendroy (Modified by Van Slyke and Hiller) *** Principle
 This is an application to urine of the blood-shorde method described on p 027 The
 urine is shake with solid eliver todate chlorde present forms insoluble sixer chlorde
 and an enjurialent amount of soluble todate. After removing insoluble material the
 odate in solution is converted to free todane which is then titrated with standard
 throculfate The method is recommended because of its simplicity and accuracy

Procedure 2st Treat 1 ml of urine with 25 ml of phosphoric tungstic acid reagent and shake with eliver lodate, exactly as described for the analysis of 1 ml of plasma or serum on p 628. After fill-ration or centrifugation treat 10 ml of the filtrate or centrifugate with sodium lodide and titrate the liberated lodine with 0 02308 N thiosulfate solution, likewise as de scribed on p 628.

CALCULATION The thiosulfate is of such strength that at the dilution of unne employed 1 ml is equivalent to 10 milliequivalents of chloride per liter of unne Ti erefore

MI 0.02308 \ throsulfate required × 10 = milhequivalents chloride per liter unine.
One milhequivalent of chloride (quals 0.085 g. of sodium chloride)

If the cloude content of the urme is so low that less than 5 ml of thosulfate are required for the titration repeat the analysis using 5 ml of urme and 25 ml of phosphore tungstic acid reagent. Shake with nodate and titrate a 10-ml portion of filtrate as before, and calculate as follows

MI 0.02308 \ thosulfate required \times 2.31 = milhequivalents chloride per liter unner.

Multiply the result by 0.085 to express m terms of g sodium chloride per liter.

Interpretation. See above

¹²¹ Harvey Arch Internal Med., 6 12 (1910)

ns See footnote 209 p 627

¹¹⁴ Reagents Lequired See method for blood chlorides p 627

FLUORINE

Determination of Fluoride (Method of Icken and Blank) ¹¹⁵ Principle Fluoride ions react quantitatively to form a complex with thorium alizarin lake, resulting in a bleaching of the color The residual intensity is read spectrophotometrically and interpolated on a reference curve derived from standard sodium fluoride solutions. The determination of fluorides in urine is preceded by the separation of phose phates and sulfates (by distillation from perchloric neid¹²¹⁸) and of chlorides (by precinitation with silver sulfation.

Procedure (a) ISOLATION OF FLUORIDES 228 Evaporate to dryness an allquot of a 24-hour specimen of urine, containing from 100 to 200 gg fluorine (usu aliy about 200 ml), hy adding it in small portions to 25 ml of a suspension of calcium hydroxide 223 in a platinum dish or crucible Dry thoroughly under an infrared lamp, then char slowly and finally ignite in o muffle oven at 600° C until all traces of ordanic matter disappear.

Cool and wet the ash with 10 ml distilled water Dissolve the ash in a minimum quantity of perchloric acld, "" keeping the dish covered with a watch glass. Rinse down the cover with a little water and transfer the solution through a funnel to a suitable distillation apparatus, e.g., a constant-temperature still "! Rinse the dish and the funnel with perchloric acld, using 20 ml for all the above mentioned operations. Add to the mixture in the still a quantity of crystalline silver sulfate equivalent to the content of childreds, previously determined on a separate allquot of the urline by one of the methods described under "Chlorides." Connect the still to a steam generator and steam distil, collecting about 200 ml of distillate.

Evaporate to dry ness and ignite the distillate as previously described, using an amount of the calcium hydroxide suspension sufficient to assure alkalin-try Dissolve the osh in perchioric acid and transfer to the still, which has been washed with hot 10 per cent NaOH and thoroughly rinsed with distilled water Add to the solution in the still 200 500 mg crystalline sliver sulfate to precipitate the remaining traces of chlorides Repeat the steam distillation and collect the distillate in a 150 ml volumetric flask to which a few drops of p nitrophenol indicator m and a few drops of 0 85 N KOIf have been odded keep the distillate aisaline at oil times by dropwise addition of 0 85 N KOIf So regulate this addition of aikaii that the distillate is neutralized as it approaches the mark.

²²⁴ Icken and Blank Anal Chem 25 1741 (1953)

²²⁴ Willard and Winter Ind Eng Chem Anal Ed 5 7 (1936)

¹¹⁷ McClure Ind Eng Chem Anal Ed 11 171 (1939)

²¹⁴ Association of Official Agricultural Clemists Official Methods of Avalysis (1950) with minor modifications.

³³⁹ Cate m hydroxude suspension Carefully slake co 56 g of low fluorine CaO with 250 ml distilled water and add slowly and with stirring 250 ml of 60 per cent HCiOs. Add a few glass beads and bot down to copous fumes of act 1 Cool add 200 ml of 10 oand bot down again. Repeat diluting and boiling down once more. Cool dilute and fifter through fritted glass fifter if precipitate of \$\fo\$0. has appeared. Pour solution with stirring into 11 of 10 per cent \(\text{\cap}\)a011 solution allow precipitate to settle \(\fo\$\text{\cap}\) on off the supernatural liquid 1 kemove sodume with sits by washing and centrifugurs 5 times 1 mally slake precipitate into suspension and make to 2 liters. I reserve in paraffixed bottles. Shake suspension well before using.

²⁰⁰ Perchloric acid Dilute 60 per cent HCiO; with 3 volumes of di tilled water and boil down to original volume. Avoil active furning Repeat and preserve in parex glass bottle.

²²¹ Huckab 13 Welel and Metler Anal Chem 19 151 (1917)
222 p-\u00e4drophenol indicator 0 5 per cent alcohojic solution

(b) DETERMINATION OF FLUORIDES Paper standard solutions of sodium fluo ride containing u, to to to to to to to g of fluoride lon, and an aliquoi of the final distillate containing from 0 to 50 mg of fluoride lon, into 50 ml of the final distingte command of the thorium alliarin reagent 114 to each flash volumetric flashs Add 10 ml of the thorium alliarin reagent 114 to each flash volumetric flashs Augustum with distilled water, stopper, shake, and allow to Nake the flasks to volume with distilled water, stopper, shake, and allow to Value the flasks to volume the light absorbance of the standards at a wave stand for 2 hours. Determine the light absorbance of the standards at a wave stand for 2 hours personness of a spectrophotometer equipped with 10 mm length of 525 m_{μ} by means of a spectrophotometer equipped with 10 mm length of 525 mg by means of a specific productive equipped with 10 mm Cores cells. Set the instrument for 100 per cent transmittance with the 500 gg. Corexcells Set the instrument of the set the instrument of the set the standard Flot the effuoride content from the reference curve

a = mL of urine taken for analysis a = mL of urine taken a is the for the spectrophotometric determination b = ml of final distillate taken for the spectrophotometric determination b if from reference a is a.

b = m or max aliquot b (from reference curve) $c = \mu g$ fluoride in aliquot $\frac{c \times 150}{a \times b} = \text{mg } \text{F}^- \text{ per liter nrine}$

Interpretation The amount of fluorine in urine is influenced by the Interpretation the diet or water supply and is therefore a enterion of fluoride content of the diet or water supply and is therefore a enterion of fluoride content of the state o fluonne expection and a crage concentration of 3 mg. F per liter of unne enamel) is prevalent an average concentration of 3 mg. F per liter of unne enamel) is prevailed as compared with a "normal" value of 1 mg per liter "bas been reported as compared with a "normal" value of 1 mg per liter "bas been reported units whose water sunniver contract the contract of the con bas been reported to those water supplies contained 0.06 and 1.36 parts per Two populations whose water supplies contained 0.06 and 1.36 parts per Two populations and of fluoride respectively (the latter by fluoridation) million (mg per liter) of fluoride levels of 0.00 ± 0.00 million (mg per man) consecutively (the latter by fluondation) three mean unner fluonde levels of 0.06 ± 0.13 and 1.12 ± 0.56 mg F showed mean blood levels of 0.4 and 1.0showed mean and blood levels of 0.4 and 4.0 µg per 100 ml respectively to per liter and respectively are the per 100 ml respectively are fluoride is eliminated not only via urine and feces but also by insentible fluoride in the persuration. No significant Fluoride is eminimated not only via urine and feces but also by insen ible and sensible perspiration to significant retention was noted when the and sensing was below 5 mg the data suggesting this to be the approximation that the suggesting this to be the approximation that the suggesting this to be the approximation that the suggesting this to be the approximation that the suggesting this to be the approximation that the suggesting this to be the approximation that the suggestion that mate limit 217

TOTAL FIXED BASE

Procedure. Measure into a large-lipped test tube (200 by 20 mm.) a sample of urine containing preferably between 10 and 25 mg. of NaCl but not more than 5 mg. of inorganic phosphorus. Add 1 ml. of approximately 4 N sulfuric acid and 0.5 ml, of concentrated nitric acid, and hoil down until white fumes appear. If the residue does not soon become colorless after this stage has been reached, cool slightly, add a few more drops of nitric acid, and continue the heating. When the remaining drop of sulfuric acid has become clear and colorless, let cool for a few minutes, and wash into a test tube which is marked at 25 ml., with four 2-ml. portions of water. Add a drop of saturated alcoholic solution of methyl red. Neutralize with powdered ammonium carhonate until the color of the indicator just begins to change, and restore the pink color by adding 4 N sulfuric acid 1 drop at a time. Heat to bolling and add more acid to restore the pink color if this is necessary. Add a 10.5 per cent solution of ferric chloride crystals in 0.2 N HCl In the proportion of 0.1 ml. for each mg. of inorganic phosphorus present, shake and run in 1 ml. of a 5 per cent solution of ammonium acetate. Add sufficient water to make the total volume 10 or 11 ml., heat again to boiling, and dilute to the 25-ml. mark with cold water. Insert a rubber stopper and mix by inverting a few times. Filter at once through a dry 9-cm. ashless paper into a dry test tube. Keep the filter nearly filled with liquid as long as possible and collect only 20 ml. Stopper the tube containing filtrate and cool. The phosphate has now been removed.

Transfer 5 ml. of filtrate to a small platinum dish, add 1 ml. of approximately 4 N sulfurle acid, and evaporate on the water bath until nearly dry. Place the dish on a metal triangle and heat cautiously at first over a microburner, gradually raising the fiame until fumes have ceased to come off. Let cool, sprinkle over the residue a little powdered ammonium carbonate, and ignite again, finally raising the fiame to its maximum and moving the triangle about until each part of the dish has been momentarily subjected to a dull red heat. When the dish has cooled, add 2 ml. of water. Agitate until the residue has dissolved, using a rubber-tipped rod to assist in solution if necessary. Transfer the contents of the dish to a large-lipped pyrex test tube, rinsing four times with 2-ml. portions of water. Determine the sulfate according to the benzidium method (see p. 949).

CALCULATION. An aliquot of one-fifth of the original urine was used for the determination and 0.02 N NaOH in the titrition. Therefore the titrition reading is equivalent to the number of ml of 0.1 N fixed base in the sample of urine used Subtract a correction of 1 per cent for the contraction of the warm solution during the filtrition stage. Results are usually expressed in terms of milliequivalents of total fixed base per litter this sobtained by dividing by 10 the number of ml of 0.1 N base per liter.

Interpretation. The total fixed base excretion, combining as it does the sodium, potassium, calcium, and magnesium exerctions, will be influenced by factors affecting any of these. In acidosis, volatile base (ammonia) plays a large part in neutralization but the fixed base exerction is also increased in varying degrees

CALCIUM

Determination of Calcium (Method of Shohl and Pedley'm): Principle. The urine is oxidized with ammonium persulfate Calcium is precipitated as oxidite

²¹⁰ Shohl and Pedley J. Biol Chem. ²⁰, 537 (1922) For references to various other procedures upon which the determination of calcium may be based, see the section on blood-calcium methods in Chapter 23, p. 644

and titrated with potassium permanganate. The method is more rapid than the gravinetric

Procedure To 100 ml of unfiltered urine ln a 250 ml Erlenmeyer flask add 5 ml of concentrated HNO, or 11, SO, and one spoonful (containing 3 to 4 g) of ammonium persulfate Insert a funnel in the flask to prevent spattering Boil and keep near the bolling point on a hot plate or over a low flame for i hour or until reduction of the persuifate is complete as evidenced by an absence of frothing when the flask is agitated. The solution at this point is pale green in color Add 10 ml of 25 per cent oxalic acid Cool to room temperature Neutralize with ammonium hydroxide, using one drop of methyl red as an Indicator Cool to room temperature If the color is now red add a few drops of ammonia to bring to intermediate color between red and yellow (pH 4 8 to 5 2) Let stand overnight Filter Whatman No 50 hardened filter paper 12 5 cm is satisfactory Wash precipitate and flask three times with distriled water,240 filling the filter two thirds full each time and allow ing to drain Break a hole in the filter paper and wash back the precipitate into the original flask, first with distilled water and then with hot dilute sul furle acld, bringing the volume to about 100 ml Add 10 ml of concentrated sulfuric acld and heat to 70° to 80° C Titrate with 0 05 N potassium per manganate taking as an end point the first color that persists 15 to 30 sec onds One mi 0 05 \ KMnO4 is equivalent to 0 0010 g Ca

Interpretation The average urmary excretion of calcium by normal adults lies between 0.1 to 0.3 g (expressed as Ca) per day. This corresponds to a to 15 millieguivalents of calcium no. Calcium excretion in the urme is dependent very largely upon the amount of calcium in the diet. From 10 to 40 per cent of the ingested calcium ordinarily is excreted by this channel the greater part being eluminated by the fees. The proportion is dependent particularly on the amount of calcium in the food. If the calcium ingestion is very high the per cent of the total excretion taking place by way of the kidneys will be low and rice tera. As excretion takes place by way of the intestine as well as by the kidneys no conclusions can be drawn from urmary analyses alone. The excretion calcium may be greatly increased in certain bone disorders as osteomalacia. In others as in rickets, the urmary excretion may be very low. For further discussion see p. 820.

The calcium content of the urine is of clinical significance in connection with the formation of certain calcium containing stones. According to Shorr **I measures designed to decrease the amount of calcium (and of phop-hate) in the urine or to increase the solubility of calcium as by promoting the urinary exerction of citrate (which forms a soluble com plex with calcium) should prove to be of value in the management of nephrolithiasis due to stones of the calcium carbonate and calcium-phosphate types.

CALCIUM AND MAGNESIUM

McGrudden's Methods Principle Unine contains magnesi im-phosphates and a small amount of iron-each of which will interfere with the accurate determinate a of its calc um cortent if proper con-litions of seed type are not martisaned dirions the

^{** --} footnote *13 p 61.

* Phort J [rol 53 .07 (1945)

precipitation In the following method the proper acidity is attained through the use of sodium acetate and hydrochloric acid, and this with slow addition of the ammonium ovalute reduces the danger of occlusion of magnesium oxalate, calcium phosphate, or ferric phosphate in the calcium oxalate precipitate.

The calcium ovalate precipitate is either ignited and weighed as CaO or determined volumetrically by titration with potassium permanganate. Magnesium is determined in the filtrate from the calcium determination after destruction of the organic matter. It is determined in the usual way by ignition of the magnesium ammonium phosphate precipitate and weighing as the pyrophosphate.

For colorimetric adaptations of these classical methods for calcium or magnesium, reference should be made to the procedures described for blood in Chapter 23.

Calcium: Gravimetric Procedure. If the urine is alkaline, make It neutral or slightly acid and filter. Take 200 ml. of the filtered urine for analysis. If it is only faintly acid to litmus paper, add 10 drops of concentrated hydrochloric acid (sp. gr. 1.20). If the urine is strongly acid, it may he made just alkaline with ammonia and then just acid with hydrochloric acid, after which the 10 drops of concentrated hydrochloric acid are added. Then add 10 ml. of 2.5 per cent oxall cacid. Run in slowly with stirring 8 ml. of 20 per cent sodium acetate. Allow to stand overnight at room temperature or shake vigorously for 10 minutes. Filter off the precipitate of calcium oxalate on a small paper, and wash free from chlorides with 0.5 per cent ammonium oxalate solution. The precipitate may then be dried, ignited to constant weight, and weighed as calcium oxide, or it may be manipulated volumetrically as described below.

Volumetric Procedure. If free from uric acid, the calcium oxalate precipitate may be washed three times with distilled water, filling the filter about two-thirds full and allowing it to drain completely before adding more. A hole is made in the paper and the calcium oxalate washed into a clean flask. The volume of the fluid is brought up to about 50 ml. and 10 ml. of concentrated sulfuric acid added. Titrate with standard potassium permanganate solution to a pink color which endures for at least a minute.

CALCULATION In the gravimetric procedure, convert the weight of CaO into terms of Ca by multiplying by 0.715. In the volumetric procedure, one mi. of 0.1 N permanganate solution is equivalent to 2.0 mg. of Ca. Calculate the daily output of calcium.

Magnesium: Gravimetric Procedure. Transfer the filtrate from the determination of calcium as above to a porcelain dish, add about 20 ml. of concentrated nitric acid, and evaporate to dryness. Heat the residue over a free flame until the ammonium salts are destroyed and the residue fuses. After cooling, take the residue up with water and a little hydrochioric acid and filter if necessary. Dilute to about 80 ml., nearly neutralize with ammonia, and cool. Add a slight excess of sodium acid phosphate and then ammonia drop by drop with constant stirring until the solution is alkaline, and then add enough more slowly with constant stirring to make the solution contain one-fourth its bulk of dilute ammonia (sp. gr. 0.96). Allow to stand overnight. Filter and wash free from chlorides with alcoholic ammonia solution (1 part alcohol, 1 part dilute ammonia, 3 parts water). The precipitate with filter paper is incinerated slowly and carefully with a good supply of air to prevent reduction, in the usual manner, and ignited and weighed as the exprohesolutare.

CALCULATION, To obtain the weight of Mg, multiply the weight of magnesium pyr phosphate by 0 2184

Interpretation. The daily exerction of magnesium by way of the urine usually amounts to between 0 05 and 0 2 g (expressed as Mg). This amount corresponds to 4 to 20 milliequivalents of magnesium ion per day. The amount depends mainly upon the dict Usually less than 50 per cent of the excreted magnesium is eliminated by the kidneys, the major portion passing out in the feces The proportion varies, however, and it is impossible to draw any conclusions from the urmary output alone There may be a retention of magnesium in certain bone disorders accompanying a loss of calcium-in osteomalacia, for example Thus the excretions of calcium and magnesium do not necessarily run parallel

Determination of Calcium in Ash of Foods or Feces.24 Ignite the material in a crucible to a white ash and dissolve the ash with the aid of a little hydrochloric acid. Bring the volume of the ash solution to 75 to 150 ml, Make just alkaline with strong ammonia added drop by drop (using litmus paper or alizarin as an indicator). Add concentrated IiCl drop by drop until just acid to litmus. Then add 10 drops of concentrated 11Cl (sp. gr. 1.20) and 10 ml. of 2.5 per cent oxalic acid. Either of two procedures may then be followed. (a) The solution is boiled until the precipitated calcium oxalate is coarsely crystalline, and then an excess of 3 per cent ammonium oxalate is slowly added to the boiling solution and the boiling continued until any further precipitate is coarsely crystalline. (If but little calcium is present, nothing will precipitate at this point and it is not necessary to add oxalate.) Or (b) the flask closed with a rubber stopper is chaken vigorously for 10 minutes. An excess of 3 per cent ammonium oxalate is then added. Cool to room temperature. Add 8 ml. of 20 per cent sodium acetate solution. (In case of ash of feces add 15 ml.) The solution may either be (a) allowed to stand overnight or (b) stoppered and vigorously shaken for 10 minutes The calcium oxalate is filtered off on a small ash-free paper and washed free from chlorides with 0.5 per cent ammonium oxalate solution. Either of two procedures may next be followed: (a) The precipitate and filter are dried, and burned in a platinum or porcelain crucible to constant weight as CaO. (b) The precipitate is washed three times with cold distilled water, as given under the method for urine and the oxalate titrated with potassium permanganate.

Magnesium is determined in the filtrate from calcum just as given above.

SODUM AND POTASSIUM

Determination of Combined Sodium and Potassium 24 From 50 to 100 ml. of urine, depending upon the specific gravity, are oxidized in a Kjeldah! flash with nitric and suifuric acids as in the Neumann procedure for total phosphorus (see p. 953). To remove the suifuric acid as completely as possible, transfer with the aid of a littie water to a platinum dish?" and evaporate to dryness over a free flame. (The alkalies are in the form of sulfate and do not volatilize.) Dissolve the residue in hot water with the aid of a little dilute hydrochloric acid. Heat to boiline and add borium chloride solution until no more precipitate forms. While still hot, add an excess of ammonio and ammonium carbonate. The harium chloride precipitates the sulfates and nart of the phosphates: the ammonia in the presence of excess harium precipitates the rest of the phosphotes, and the carbonate precipitates the calcium and most of the magnesium, as well as the excess horium, Filter and wash the precipitate well with hot water containing a few drops of ammonia. Evanorate the filtrate and washings to dryness and heat the residue to duil redness for a moment. Redissolve in water and treat again with ammonia and ammonium carbonate to remove any remaining alkaline earth metals. Filter and wash as before. Transfer the filtrate and washings to a weighed platinum dish, add a few drops of hydrochloric acid, and evaporate to dryness. Heat the residue dentiv to remove ommonium salts and then to dull redness for a moment Desiccate and weigh, Reheat to constant weight which represents the combined chlorides of sodium and potassium. The reasents used in the determination must be tested and found free from alkali metals or a correction made for the all ali metals present in the reasents used. The sodium is determined by difference after notassium has been estimated by the method given below.

Patassium. Dissolve the alkali chlorides from the preceding determination in a little water and add a slight excess of 10 per cent platinic chloride over thot necessary to precipitate all of the alkali present calculated as sodium chloride (ahout 17 ml. being required for each gram of sodium chloride). Evaporate to a syrupy consistency on the water hath and add about 50 ml. of 80 per cent alcohol. Stir occasionally for a few minutes. This operation must be carried out in the absence of ammonia vapors. Fliter through a weighed Gooch crucible, washing the precipitate with 80 per cent alcohol first thoroughly by decantation and then on the filter, for some time after the filtrate is colories. Dry of 10° to 15° C. and weigh.

CALCULATION. Multiply the weight of potassium plations chloride by 0 1608 to obtain the amount of K prescot Express as KCl by using instead of this factor the factor 0 3067. Subtract from the weight of total alkali chlorides as determined in the preceding method, the weight of potassium chloride as calculated and obtain the amount of sodium chloride present. To convert sodium chloride into sodium, multiply by 0 3034. To express the sodium (or potassium) content in terms of miliequivalents, dayide the weight of NaCl (or KCl) in gramps by 0 30845 for 0 07455).

Interpretation. The average alkali everetion of an adult on a mived diet is about 1 to 3 g. of potassium expressed as K (40 to 65 milliequivalents), and 30 to 5 g. of sodium expressed as Na (130 to 200 milliequivalents). The ratio of Na to K is thus about 5:3. Both the ratio and the absolute amounts of these elements exercted are, however, largely dependent upon the salt content of the diet. The urine during fasting contains more potassium than sodium salts, because of the noningestion of sodium chloride and the accompanying destruction of potassium-containing body tissues. The excretion of the bases, particularly K, may be increased in fevers and in acidosis.

IRON

Methods of Fivehjem and Kennedy. The urine is ashed, the ash dissolved, and the iron present determined colorimetrically as thlocyanate.

Procedure. Evaporate and ash 100 ml. of urine and carry out iron determination according to Elvehjem or Kennedy (see p. 656).

IODINE

Method of Von Fellenberg-312 Principle. For discussion of the principle, 500 the original articles 'McCullaghan' has suggested a simpler procedure for jodine determinations in blood and other materials Chaney's method 317 for protein-bound jodine is described on p. 661.

Procedure. To 10 to 40 ml. of urine add I to 3 mL of a saturated solution of lodine-poor K₂CO₂ and evaporate in a low iron dish (about 10 cm. diameter). Heat gently without igniting, molsten with water, and Ignite, not bringing the bottom of the dish to redness. Extract the charted mass with a little water and filter. Ignite the paper and residue, then add the filtrate (only falloty) yellow) and a few drops of 10 per cent NaNO₃, and complete ignition. The residue should he pure white. Extract Iour times with 2 to 3 ml. of alcohol. Evaporate the alcoholic solution in a gold or platinum dish (about 6 cm. diameter) on a water bath to dryness, after adding a few drops of saturated K₃CO₃ solution. Ignite gently. Extract again with 95 per cent alcohol and evaporate and Ignite, this time adding no K₃CO₃. The bottom of the dish must not turn red. Dissolve the residue in 0 3 ml. of water and titrate Iodine according to the method of McCOllaghtim or according to the procedure of v. Fellenberg. The amount of Iodine eliminated In 24 hours may be from 10 to 200 ad.

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For the determination of urinary 17-ketosteroids, see Chapter 26 For the determination of vorious vitamins in urine, see Chapter 35 See also

The method given is a modification by Lunde Biochem. Z., 193, 94 (1929).
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¹⁰ Chaney Anal. Chem., 21, 933 (1950)

von Fellenberg Prob. Physiol 25, 176 (1926) Biochem. Z., 224, 170 (1930) Lunde
 Closs and Pedersen Biochem. Z., 284, 264 (1925)

the references to methods for blood analysis (Chapter 23) since many of these are equally suitable for urine

TESTS FOR KIDNEY EFFICIENCY

I Blood-Urea Clearance (Method of Moller, McIntosh, and Van Sijse) absorbed in the tubules The blood urea clearance test is a measure of the efficiency with which the kidney everetes area Results are expressed either in terms of the volume of blood in ml which is completely cleared of urea per minute, or more commonly as the percentage of the average normal value which this volume represents. The necessary data include the concentration of urea in blood and urine and the volume of urine exercted in unit time.

Procedure The patient is permitted to eat a moderate meal (usually breakfast) without coffee \ignorous exercise is avoided and after breakfast the patient is kept at rest during the 2 hour test period. At the beginning of the test period, the biadder is emptied and the urine discarded. The patient is given a glass of water to drink, and the urine is collected in two succeeding periods of one hour each. At the beginning of the second period, the patient is given another glass of water, and a sample of blood is drawn for uren determination. At the close of the test, the volumes of urine passed during each period are accurately measured in graduated cylinders, and the urea content of each determined separately.

The two collection periods need not be exactly one hour long, but may be longer if convenient. It is important, however, that the time of the period he accurately known, to the nearest minute, and that the entire volume of urine formed during this period he obtained, since only in this way can the volume of urine flow per minute he calculated. The two periods serve as checks on one another, if results calculated on the basis of each separate period differ significantly, error is indicated. If the urine volume is less than 25 ml per hour, the sample is discarded

Calculation The maximum clearance (C_w) indicates the maximum efficiency of urea exerction with high urine volumes and is calculated if the urine volume observed in an adult, or if the corrected volume $V \times \frac{173}{\text{sq m}}$ surface area. In a child, exceeds

2 ml per minute. The standard clearance (C_s), or the efficiency with which the kidneys excrete urea when the urine volume is at the average normal level of 1 ml per minute, is calculated if the urine volume, corrected in the case of a child is less than 2 ml per minute. Both clearances are best calculated in percentage of the average normal C_s or C_m

Percentage of average normal
$$C_n = \frac{100~U1}{70B} = 1~33~\frac{U1}{B}$$

Percentage of average normal $C_n = \frac{100~U~\sqrt{1}}{54B} = 1~85~\frac{U~\sqrt{V}}{B}$

U = Ures concentration of urine (mg urea > per 100 ml)

B = Urea concentration of blood (mg urea \(\sigma \) per 100 ml)
 V = ml urine exercted per minute

²⁰ Möller Meintosh and Van Siyke J Clin Invest 6 42" (1929) See also Van Siyke and Cope Proc Soc Exptl Biol Med 29 1169 (1932)

The following table is convenient in calculating standard clearance. It gives values of \sqrt{V}

V ml per minule	$\sqrt{\overline{v}}$	I' ml per minute	$\sqrt{\overline{v}}$	V ml per minule	\sqrt{V}	V ml per minule	$\sqrt{\vec{v}}$
0 2	0 45	0 7	0 84	1 2	1 10	1 7	1 30
0 3	0 55	0 8	0 89	1 3	1 14	1 8	1 34
0 4	0 63	0 9	0 95	1 4	1 18	1 9	1 38
0 5	0 71	1 0	1 00	1 5	1 23	2 0	1 42
0 6	0 78	1 1	1 05	1 6	1 27	2 1	1 45

Interpretation. In patients with diminishing renal function, the blood-urea clearance shows evidence of diminution sconer than does the blood-creatinine content, the blood urea content considered without relation to urea excretion, or the phenolsulfonephthalein excretion. The blood-urea clearance usually falls below 50 per cent of normal values hefore any of the other three show abnormality. Only after the blood-urea clearance indicates less than 20 per cent of normal renal function are all values for blood urea, creatinine, and phenolsulfonephthalein found outside the limits of normal variation. The maximum clearance is normally about 40 per cent greater than the standard clearance, the mean values being 75 ml (variations 64 to 99 ml) of blood per minute for the maximum and 54 ml (variations of 40 to 68 ml) for the standard. The method is based on the view that with abundant urner the urea excretion per minute equals the urea contained in a constant volume of blood.

Phenoisuljonephthalein Test Principle This test for renal function was devised by Rowntree and Geraghty. It depends upon the injection into the tissues of a dyestuff which is eliminated rapidly by the normal kidneys, and can be easily estimated quantitatively in the urine.

This dyestuff, phenoisulfonephthalem, is nonirritative to the body either when taken by mouth or when injected into the tissues so that it does no harm to an already weakened kidney

The patient upon whom the test is to be performed is given 300 to 400 ml of water 20 to 30 minutes previously, in order to assure a free flow of urine. Just before the start of the test, the bladder is empired and the urine discarded.

Procedure One mi of a solution containing 6 mg of phenolsulfonephthaleinine per mi is injected intramuscularly an the lumbar region, the time of injection being noted. The patient is then catheterized and the urine as it forms thereafter allowed to drop into a beaker containing 2 drops of 25 per cent haoli! The appearance of a red color in the alkalinized urine indicates beginning excretion of the drug, the normal time being within 5 to 10 mioutes after its injection. Urine is now collected in one-hour samples in

¹¹⁸ This solution is prepared by adding 0.5 g. of phenolvalionephthalein and 0.54 ml of 2 N xoll it co acough 0.75 per cent had Castion to make 100 ml. This gives the more solution or acid salt which is shightly irritant should be injected. It is necessary 70 add two to three drops more 2 N xoll which changes the color to a Bordeaux red This preparation is noniritant Suitable preparations of the dye in sterile ampals may be obtained from pharmaceutical supply houses.

patients with obstruction to the flow of urine from the bladder, the retention catheter is stoppered and the urine drawn off at the end of each hour. Other patients may simply be allowed to urinate at the hourly periods.

To each hour sample of urine is added 25 per cent NaOH, drop by drop, until the maximum intensity of color appears. This color will remain constant for an indefinite period of time. Each sample is then placed in a 1000-ml, volumetric flask and diluted to the mark with distilled water.

Compare the color intensity of each sample either colorimetrically or photometrically against a standard. To prepare the standard, place a sufficient amount of the phenolsulphonephthalein solution to contain 3 mg, of the dye (i.e., one-half of the amount administered) in a beaker, dilute with a little water, and add 25 per cent NaOH dropwise to maximum color intensity. Transfer quantitatively to a 1000-mi. volumetric flask, dilute to the mark with water, and mix.

For colorimetric comparison, match the unknowns against the standard in the usual way. It is convenient to set the standard at 10 mm. For photometric measurement, determine the densities of standard and unknown in a photometer at 520 mm, setting the photometer to zero density with water.

CALCULATION For colorimetric measurement

Reading of Standard Reading of Uaknown × 50 = percent of administered dye io sample

For photometric measurement

Density of Uoknowo × 50 = per cent of administered dyom sample

Interpretation. The amount of the drug eliminated normally is 40 to 60 per cent during the first hour and 20 to 25 per cent during the second hour, or a total of 60 to 85 per cent for two hours. The amount of the drug excreted has been found to be independent of the quantity of urne obtained. Ordinarily, two one-hour samples are sufficient; in case of delayed excretion the collection of hourly samples may be continued until practically all of the drug has been recovered in the urine

If it is desired to test the function of each hidney separately, ureteral catheterization must be resorted to, the experiment otherwise being performed as above described.

The phenolsulfonephthalem test may be used to indicate the amount of derangement in quantitative functional disturbance of the kidneys, as in chronic interstitial and chronic parenchymatous nephritus or uremia.

It is claimed that the rate of exerction of phenolsulfonephthalein is affected by certain extrarenal factors; namely, the albumin and hydrogen-ion concentrations of the blood.

 Mosenthal Test for Kidney Function:²¹¹ Principle. The patient under examination is placed for a day on a more or less definite diet.²¹² The urine is collected

^{33.} Mosenthal Boston Med. Surg. J., 17a, 245 (1914), Ohio State Med. J., 18, 345 (1922), 36 A diet suitable to ordinary hospital conditions is given by Kahn Functional Diagnosis, p. 260, New York, W. F. Prior Co., 1920. It is not essentful, as was formerly believed to prescribe a diet abundant in dimertic foods or beverages, since ordinary foods contain sufficient diruction materials for the proper carrying out of the test. In private practice it is only necessary that the primite set three full meals a day and record the approximate quantities—1 cup of coffee, 2 shees of toxic, 2 tables/poorpuls/of ordened, etc.

in six two-hour periods during the day and one 12-hour night period. These unne specimens are analyzed for volume, specific gravity, total nitrogen, and chlorides

Procedure. On the day of the test have the patient empty the bladder at 8.A.M. and start the diet for the day which, if desired, may contain approximately 13 to 14 g. of nitrogen, 8 to 9 g. of salt, 1700 to 1800 ml. of fluid, and considerable purine material in meat, soup, tea, and coffee, "" No solid food nor fluid of any kind must be taken between meals and especial care must be observed that nothing is eaten or drunk after the evening meal. The meals should start at 8.A.M., 12 noon, and 5 P.M.

Collect the urine punctually at the end of every 2-hour period until 8 P.M., and place in separate bottles. Collect the night urine from 8 P.M. to 8 A.M. of the following day in another bottle. Measure the volume of each specimen of urine and determine in each case the specific gravity, total nitrogen, and total chlorides.

Interpretation. The test is of particular value apparently as giving earlier indications of diminished kidney efficiency than is true of some other tests used. It is sometimes difficult to interpret the results obtained in terms of renal involvement because of the influence of possible extra-renal factors. In general, however, the normal response is one in which the specific gravity figures vary at least 9 points (less if too little water is taken) from the highest to the lowest and the volume of the night urine is 400 ml. ²³⁴ or less if the percentage of nitrogen and sodium chloride in the night urine or in the highest of any of the day specimens is 1 per cent, a normal condition is indicated. Values under 1 per cent, however, may or may not be abnormal

Time of Day	Urine		Sodium Chloride		Astrogen	
1 tme of Day	MI	Sp gr	Per Cent	g	Per Cent	G
8-10 10-12 12-2 2-4 1-6	153 156 191 260 114	1 016 1 019 1 012 1 014 1 020	1 32 1 25 0 61 0 77 0 93	2 02 1 95 1 24 2 00 1 13	0 89 0 74 0 59 0 56 0 95	1 26 1 15 1 14 1.46 1.98
6-8 Total day Night, 8-8	238 1115 375	1 020	0 43	9 36	0 52	1 235 7 32
Total, 24 hours Intake	1499 1769	1 020	0 63	2 36 11 72 8 50	1 23	11 93 13 40
Balance	+270	1		-3 22		+1.47

When kidney function becomes involved, the first signs are usually demonstrated in the night urine. The quantity becomes increased and

¹⁴ Ger footpote 252

¹⁶ This represents the usual normal limit. Volumes in excess of 750 ml are distinctly abnormal, whereas volumes between 400 and 750 ml are of doubtful significance.

the specific gravity and the nitrogen concentration are lowered. One or all of these changes from the normal may occur. In severe cases of chronic nephritis an advanced degree of functional inadequacy of the kidney is indicated by a markedly fixed and low specific gravity, a diminished output of both salt and nitrogen, a tendency to total polyuna, and a night urine showing an increased volume, low specific gravity, and low concentration of nitrogen. Such functional pictures are, however, not confined to nephritis. They are found frequently in many other conditions pyelitis, cystitis, hypertrophied prostate, marked anemia, pyelonephritis, polycystic kidney, and diabetes insipidus. The table above taken from Mosenthal shows the response of a normal individual

4. Other Methods. Inulin is filtered through the glomeruli and notther excreted nor reabsorbed by the tabules This is the basis for an inulin clearance test for renal (glomerular) function 255

The p-aminohippune and (PAH) clearance test is employed as an index of rend blood flow since this compound is filtered through the glomeruli and exerted via the tubules *st

Other hidney function tests based on dye exerction rates or on concentration and dilution efficiencies have been proposed

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²³⁴ Miller Mying and Bulin J Clin Invest 19 89 (1940)
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32

Isotopes

The chemical properties of the several elements depend upon the number and configuration of the unit negative charges, or electrons, which surround the nuclei of their ntoms. The number of such planetary electrons in the neutral atom of any given element is equal to the number of protons, each having a unit positive charge, in the nucleus of that atom. An element is characterized by a fixed and definite atomic number of protons in the nuclei of its atoms, balanced by an equal number of planetary electrons.

All atomic nuclei, with the single exception of the nucleus of ordinary hydrogen, contain one or more neutrons in addition to protons. The mass number of the neutron (1 00890) is almost identical with that of the proton (1 00813), based upon the arbitrary designation of 16 for the mass number of the commonest variety of oxygen. The number of neutrons in the nuclei of the atoms of a given element is not necessarily fived, but may vary within narrow limits. Therefore the atoms of most elements contain nuclei with different numbers of neutrons, and consequently of different mass numbers. Each such atomic species, with a specific mass number as well as a specific atomic number, is called an asslone.

The complete symbol for any isotope includes the chemical symbol of the element preceded by the atomic number written as a subscript and followed by the mass number written as a superscript. Thus ordinary oxygen would be written 60¹⁴, ordinary carbon as 60¹⁴, and ordinary hydrogen as 1H. These elements also have less abundantisotopes such as 60¹⁴, 60¹⁸, 61¹⁸, 1H, and 1H. Since the atomic number is implicit in the chemical symbol, it is more common practice to omit the subscript and simply write 0¹⁴, Ci¹³, H, Or, Ou, Ci¹³, H, or H for the isotopes mentioned in the previous sentences An equally logical but less often used system is to give the two numbers, omitting the chemical symbol.

Most elements exist in nature as mixtures of isotopes. The table on p 971 lists the isotopes of the elements of major physiological significance. The proportions of the several isotopes in a naturally occurring sample of an element are, as a general rule constant regardless of the source of the material. For example, the percentage of O¹⁷ is the same in atmospheric oxygen and in oxygen obtained by decomposition of ocean water or of silicate or carbonate rocks. Most of the naturally occurring isotopes listed in the table are stable, which means that they do not exhibit radio-activity as a result of soontaneous nuclear deeas.

For any given element with its fixed number of protons there custs only a limited number of possible isotopes early with its specific number of neutrons Of the a isotopes still fewer possess stable index. Among certain elements usually those containing an odd number of protons only one of the possible isotopes is stable sodium phosphorus, and iodine are examples of such elements with only one stable isotope. Elements whose nuclei contain an even number of protons are likely to have more than one stable isotope. Fin for example has 10 As atomic nuclei in crease in size and complexity, the proportion of neutrons required to maintain stability also increases. From the table on p. 971 we can see that iron (atomic number 26) requires 30 neutrons to stabilize its most common isotope, the neutron proton ratio is 11. It oldine (intomic number 33) has for its stable isotope a neutron proton ratio of 1.40 while the most complex of all stable elements is bit muth (atomic number 83), who e 126 neutrons give it a neutron proton ratio of 1.52.

All nuclei more complex than that of bismuth and many that are less complex are unstable and undersy spontaneous changes in neutron proton composition Such changes constitute radioactivity and may occur

in one or more of three ways

Upha perticles each convising of 2 protons and 2 neutrons may be emitted resulting in a loss of 4 in mass number and of 2 in atomic number. Beta particles which are high peed electrons may be emitted cauling a gain of 1 in atomic number by the change of a neutron into a proton, and no change in mass number.

Gamma rays which are electromagnetic radiations comparable to x rays but more energetic and penetrating may be emitted and cause no change in mas or atomic number. The production of γ rays and the high velocities at which α and β particles travel are evidence of the release of nuclear energy previously stored in the bonds between protons and neutrons. Such nuclear energy is of an order of magnitude sufficient to permit its evaluation in terms of mass according to the relationship demon strated by Einstein that $0 \times 10^{\circ}$ ergs of energy are equivalent to 1 gram of mass

When these energetic radiations impinge upon surrounding atoms of molecules they not only produce chemical changes by altering the configuration of planetary electrons but they also cause changes in nuclear structure transmuting one element into another. The cyclotron and the atomic pile are devices for the production of high-energy radiations in high concentrations. By their use radioactive isotopes can be prepared in useful quantities. Radioactive isotopes are now known for every chemical element. Both types of isotopes stable and radioactive have been useful in biological investigation. The two types require different procedures for their utilization and have different applicabilities. Each type will be considered separately.

RADIOACTIVE ISOTOPES

Radioactive isotopes differ with regard to availability half life and type of radiation emitted 'wailat lity now appears to be largely a matter of technical development. The type of radiation emitted (kind and intensity) influences the sensitivity and methods of measurement. The half life of a radioactive element is the time required for one-half of the atoms to disintegrate. Isotopes with a very sbort or very long half-life are of less value than those with a half-life of the order of weeks or months. A short half-life means that too great a proportion of the original material will have lost its activity (and original chemical nature) during the time required for transport after preparation to the place of use, and for the manipulative details of an experiment. A very long half-life means that a high proportion of the isotope must be incorporated into a compound if accurate radioactivity measurements are to be made.

In contrast to the ordinary radioactive half life defined in the preceding paragraph, we sometimes speak of the biological half life, which is the time taken for a radioactive element within the hody of a given organism to lose half of its activity. Elements which are firmly fixed in the hody, such as plutonium in hone, have a biological balf life comparable to the ordinary half life. Excretion of other radioactive elements causes a loss in total hody radioactivity frequently more significant than the loss by radioactive decay.

Measurement of Radioactivity. Most methods for the measurement of radioactivity depend upon the fact that energetic α or β particles cau convert atoms or molecules of a gas to positive ions by displacing electrons from them Each positive ion so formed is paired either with the freo detached electron or with a negative ion formed by the union of the electron with another neutral atom or molecule. The number of such ion pairs formed per centimeter of the path of a particle is called its specific ionization. Gases can also be ionized by γ rays, which detach high velocity electrous from atoms or molecules in their path. These secondary electrons in turn produce ion pairs

The ionizotion chamber can be used for the measurement of any type of ionizing radiation, but is particularly adaptable to a particle counting In its simplest form, an ionization chamber consists of two parallel conducting plates, usually about 1 cm apart, mounted in a chamber containing a suitable gas, with provision for maintaining a potential difference across the plates and for measuring the current flow through the circuit The radioactive material can be placed on the lower plate or alternatively the a particles may enter the chamber through a thin window of mica No current flows unless ionization of the gas is produced, when positive and negative ions will travel towards the oppositely charged plates Within the range of voltage across the plates suitable for the individual instrument (usually 100 to 200 volts), every ion formed reaches the plates and causes an electrical impulse. The instrument is designed to take advantage of the high specific ionization of a particles and to minimize the disadvantage of their short range. The ionizing chamber is usually used in connection with an amplifier and a mechanical counter, or a counting rate meter A scalar is a device used with mechanical counters to cut down, by a predetermined factor, the number of electrical impulses recorded A pulse analyzer or "kicksorter" differs from a scaler in that it transmits only stronger impulses, chiminating the weak effects of 8 and 7 radiation, and may also assort the transmitted impulses

according to strength and allot them to different counting circuits. In counting mixed radiations α particles may be screened out by a thin sheet of aluminum, and both α and β particles by a thin sheet of lead

In proportional counters the applied voltage is increased to a range (500 to 800 volts) where the electrons inherated in the primary ionization are given sufficient velocity to cause secondary ionization with formation of more electrons, which in turn are speeded up in the high-potential field, and so on This results in a self amphification of impulses known as the Townsend cascade, or Tounsend avalanche. The degree of such self

amplification depends upon the applied voltage

The Geiger-Muller counter utilizes a still higher voltage (800 to 1500 volts) with a greater degree of self-amplification and only minimal variation with small changes in applied voltage. The cascades produced under these conditions are likely to continue for some time after the entry of the initiating ionizing radiation, and so may merge with the impulses from subsequent rays or particles. To resolve individual impulses, some form of quenching is necessary, which may be internal by the use of poly atomic organic gases, or external by the use of resistances. An adequately quenched Geiger-Muller counter will resolve individual particles at rates less than 5000 per second. The probability of two or more impulse-countring during an interval too short for resolution can be calculated—the coincidence correction—for a given tube and a given counting rate Geiger-Muller counters have been designed for all types of ionizing rations, but are most suitable for β particles and γ rays.

Scintillation counters utilize the flashes of light produced, chiefly by α particles, upon impact of the radiation upon zine sulfide and certain other substances. The photomultiplier tube is used in counting circuits based on this principle Still other types of counters are actuated by the change in electrical conductivity of certain crystals when exposed to

ionizing radiations

The activity of any radioactive source can be expressed in curies. The curie was originally defined as the quantity of radion in radioactive equilibrium with one gram of radium. It is now defined by the approximate equivalent, 3.7 × 10% dismitegrations per second. A millicurie, therefore, desembes a radioactive source with 3.7 × 10%, and a microcurie such a source with 3.7 × 10% dismitegrations per second. Another unit, the rutherford, defines the quantity of radioactive material that deeps, at a rate of 10% dismitegrations per second. Note that these units refer to disintegrations, and not to particles or ray semitted. From measurements of radiation per unit time, expression of the results in curies or rutherfords requires a calculation of disintegrations per unit time utilizing the decay scheme (which may be quite complex) of the radioactive element in question. By measuring the ionization produced by any 7 ray emitter, its source stringth can be expressed in reentgens per hour at one meter (see p. 976) without knowledge of the disintegration scheme.

Radioautographj is a method for detecting and measuring radioactive substances by the effect of their radiations upon the silver bromde of photographic emulsions. The action of light and of ionizing radiations upon the emulsion is comparable in that electrons are transferred from bromide ions to silver ions, with formation of metallic silver—the latent image—around which the silver produced by further reduction in the developing process precipitates A radioautograph is therefore produced by placing the radioactive material in close contact with a photographic emulsion. Light must of course be excluded, and direct chemical action of

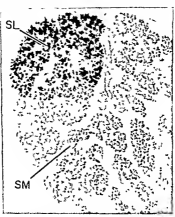


Fig. 258 Radio-Autographs of Sublingual (SL) and Submaxillary (SM) Glands of a Rat Given an Injection of Chelabeled Bicarbonate Immediately after Birth

The dark area indicates the presence of C¹-labeled substances as seen in an animal sacrificed 4 hours after injection.¹⁴

Courtesy, R. C. Greukeh and C P. Leblond,

the specimen upon the photographic emulsion may be obviated by interposing a thin layer of celloidm. Following adequate exposure, the photographic emulsion is developed, fixed, washed, and dried in the manner eustomary for ordinary photographic negatives. In bistological studies, it is advantageous to have stained section and radioautograph superimposed for observation. Several techniques for this purpose have been described. Figure 25S shows by radioautographs the comparative secre-

¹ Gross, Bogoroch, Nadler, and Leblond: Am. J. Roentgenol, Radium Therapy, 65, 420 (1951).

¹⁶ The sublingual gland (darker area) shows an intense reaction given by the nuccous material present in the vicinar cells as well as the lumn of the ducts. The elaboration and release of the sublingual mucus is extremely rapid. The submanillary secretion is considerably slower.

tory activity of the sublingual and the suhmaxillary glands in newhorn

Biological Effects of Ionizing Radiations. The unit in which exposure to x rays or to y rays is measured is the roentgen The roentgen (r) is defined as the quantity of x rays or y rays which will produce, by ionization, I electrostatic unit of electricity of either sign in 1 ml of dry air measured under standard conditions of temperature and pressure A more convenient equivalent definition for the roentgen is the amount of x-rays or y rays which must be absorbed by 1 g of air to cause it to gain 83 8 ergs of energy

To include other types of ionizing radiations, such as α and β particles, the roentgen equitalent physical or rep is defined as the quantity of ionizing radiation of any kind which, if absorbed by the soft tissues of the body, causes un energy gain per gram of tissue equivalent to that caused by

1 r of x-rays or γ rays

Since the absorption of 1 rep of one type of radiation might have a biological effect quite different quantitatively from that of 1 rep of mnother type, the roenigen equivalent man or rem is defined as the quantity of radintion of any type which produces the same amount of damage in man upon absorption as i r of x rays or γ rays. For x rays, γ rays, and β particles, 1 rep equals approximately 1 rem, but for α particles, 1 rep equals 10 to 20 rem

The combined experience and experiments of medical radiologists and of the physicists charged with the protection of workers handling radioactive materials has led to the tentative conclusion that the total permissible dose of all kinds of ionizing radiation is 0 3 rem per week over the whole body In defining this dose it is understood that very concentrated whole body exposure for short times is dangerous even within the limits of the permitted amount per week Larger doses up to 15 rem per week, may be permitted if exposure is limited to hands and forearms

Neutrons are emitted at high velocity from atomic explosions and in nuclear reactors Such "fast neutrons" possess high energies but limited powers of penetration When fast neutrons interact with tissue, they are first slowed down by collisions with nuclei predominantly those of hydrogen, which are thereby detached from their molecules and stripped of orbital electrons in this manner fast neutrons are actually ionizing radiations When the velocity of the neutrons is cut down by repeated collisions, the neutrons may be captured by ordinary atomic nuclei, bringing about nuclear transformations \cutron capture may result in the ejection from the nucleus of energetic protons, or in the emission of 7 rays accompanying the formation of deuterium in the case of by drogen, and in other instances the formation of radioactive isotopes. All of these nuclear transformations are further sources of ionizing radiation, which are spoken of collectively as induced radioactivity . certs

The detached electron can also react with a water molecule, yielding a hydroxyl ion and a hydrogen atom

The ions react together to form water molecules again. Some of the hydrogen atoms unite to form hydrogen molecules. Some of the free hydroxyl radicals unite to form hydrogen peroxide A hydrogen atom may unite with a hydroxyl radical to form water. Thus radiation of pure water forms hydrogen and hydrogen peroxide.

If the water contains dissolved oxygen, it will combine with hydrogen atoms, increasing the yield of hydrogen perovide, and forming per hydroxyl radicals

$$H + O_2 \rightarrow HO_2$$

Among the substances which have been shown to diminish the destructive effect of ionizing radiations upon solutions of enzymes or upon hving cells, cysteme and other substances containing the sulfhydryl group are noteworthy. The sulfhydryl group can reduce hydroxyl radicals, hydrogen peroxide, and perhydroxyl radicals, thus removing from aqueous solutions these products of ionizing radiation. The other product, hydrogen atoms, is removed indirectly through the formation of perhydroxyl radicals (in the presence of dissolved oxygen, see equation above)

The lethal effects of radiation result chiefly from the destruction of the more susceptible cells (all types of blood and epitbelial cells) and the failure to form new cells to replace them Death after a single heavy exposure is usually delayed weeks or months, during which time blood cell counts fall and ulcerations of epithelia develop. The symptoms which appear promptly after beavy irradiation—the initial radiation response or radiation sickness-do not necessarily indicate a lethal exposure Gastrointestinal upset and lowered blood pressure may occur even after heavy 1-ray dosage within the therapeutic range, particularly if applied to the abdomen Possible mechanisms to explain this prompt reaction are the liberation of VDM (to be discussed later) and direct effects of radiation on the autonomic nervous system. The damaging effects of ionizing radiations upon tissues can be demonstrated instologically 4 The clearest results are obtained in specimens prepared by freezing-drying, then cut and mounted dry Vacuoles appear within the cell nuclei of tissues which have been subjected to radiation. In cells such as lymphocytes, which are sensitive to radiation, the vacuoles distend and eventually rupture the nucleus. In more resistant cells, the vacuoles formed are smaller, and are later extruded through the nuclear membrane without necessarily destroying the cell

The well-recognized genetic effect of sublethal exposures to ionizing radiations is an increase in the rate of occurrence of mutations. It has not been definitely extablished whether this effect is chemical (upon the

Swallow J Chem. Soc., 1334 (1952)
 Warren Holt and Sommers Am. J Clin Path 22, 411 (1952)

nucleoprotein molecule) or mechanical (upon the chromo-ome) The genetic effect of radiations has been shown to be cumulative, depending more upon the total radiation received by the experimental plant or animal than upon the intensity of the radiation. Data relating amount of radiation to genetic effects in man are not available. Sterility can be produced by ionizing radiations in sublethal doses.

The amount of energy expended in producing the lethal effects of ionizing radiation in living cells is very small, less than one-millionth of the cell's normal daily expenditure of energy in metabolic processes. It is therefore exident that damage to key substances, such as enzymes or nucleic acids, by ionization must be postulated to explain the destructive action of relatively minute amounts of energy. Sulfin, dryl enzymes are particularly susceptible to mactivation by radiation, even in vitro. One chemical mechanism by which tissue damage from radiation is known to take place is the formation of peroxides. Two-thirds of the total biological effect of exposure of Paramecium aurelia to x rays could be attributed to peroxide formation in the medium, since it could be abolished by catalase and could be duplicated by adding H₂O₂ to the medium without radiation.

Ionizing radiation also brings about the release within the circulating blood of a toxic substance which causes severe peripheral vasodepression. The behavior of this substance is like that of a similar substance betweeted in traumatic shock. This substance was first designated simply as VDM (vasodepressor material) but was later identified as ferritin. By several criteria based upon physiological action, ferritin and the VDM of irradiated animals are identical. A vasoexitor material which has not been identified chemically, has been reported in situations of reversible shock and in animals that survived into the second wick following irradiation.

Radioactive Tracers. Physiological chemists have repeatedly sought a means of placing an identifying tag or label upon a particular sample of a foodstuff, drug, or intermediate and following it through its metabolic transformations in a living organism. Knoop used such a tracer in his studies of fatty-acid oxidation when he substituted the phenyl group on the omega carbon atom of fatty acids. Eladic acid, the trans isomer of loice acid, has been similarly used. In both these instances the pathway of the tracer substance was found to be not identical in all metabolic sequences with that of the natural substance under my estigation.

The use of tracer substances identical with the corresponding natural substances except for one or more radioactive isotopic atoms in the molecule has obtained this difficulty. The tabsed atoms can be identified by their radioactivity, whatever the compounds may be into which they are introduced by metabolic activities. Such tracer substances can be prepared by ordinary chemical synthesis or in some instances more efficiently by biosynthesis using as starting inaterial radioactive isotopes obtained by the use of the cyclotron or the nuclear reactor. The fundamental

postulate in metaholic studies with radioactive tracers is that the chemical changes undergone by the radioactively tagged molecule during its metaholism are the same as those if the corresponding untagged natural substance. This postulate is valid if, and only if, two important experimental conditions are maintained (a) radioactivity must be kept at a level low enough not to cause disturbance of cell metaholism, as for example by inactivation of enzymes, and (b) the tracer substance must be given in amounts not large enough to exceed physiological concentrations in the organism. The latter condition can be purposely waived if the study involves the effects of unusual concentrations or unphysiological substances.

Hevesy⁸ made the first hological application of radioactive tracers He followed the absorption and distribution of lead in plants with thorium B (in modern terms, Ph²¹²), a naturally occurring radioactive isotope of lead The choice of available radioactive isotopes at that time was, of course, extremely limited Currently the nuclear reactor and the cycloron make available a large number of radioactive isotopes of elements significant in hiology. The radioactive isotopes most useful in present metaholic studies are Ci⁸, H², Pr², Na²⁴, I¹²³, Fe²⁴, Fe²⁵, Si²⁵, and Co¹⁶ Procurement of radioactive isotopes for experimental work requires the authorization of the Atomic Energy Commission. No useful radioactive isotopes exist for two very important elements, oxygen and nitrogen Tracer work with these elements must therefore he done with the stable isotopes, O¹⁵ and N¹⁵.

Although samples of pure radioactive isotopes can he isolated (with considerable difficulty), the material used for preparations of tracer substances is almost invariably a mature of the desired radioactive isotope with stable isotopes of the same element. The specific activity of a sample is defined as the number of atoms of the specified radioactive isotope divided by the total number of atoms of the same element. (Unfortunately, the term specific activity is also used to mean curies per gram of the element specified. The reader must distinguish these two different quantities by the units used.)

An important consideration enters here into the interpretation of results obtained with labeled organic molecules. If the isotopic element is in a labile position—i.e., if it can enter readily into exchange reactions with the solvent or with other elements in the molecule—interpretation is obscured. For example, an organic acid labeled with deuterium in the iomizable portion of the carbox1 grapp may exchange with the nonsotop c hydrogen of water present to render the labeling valueless as a means of identifying the compound

$R COOD + H_2O \rightleftharpoons R COOH + D_2O$

Thus a stably hound isotope is in more value in this connection than a lable one, and due consideration must be given to this fact. It should be pointed out that some positions in organic molecules are stable in the but may become lable in the Thus the α amino introgen of an immo

¹ lleves) Buchem, J 17 439 (19.3)

acid may be stable, but may become detached by metabolic processes so that the remainder of the molecule is no longer identifiable in terms of isotope content:

R CHN14H2 COOH - R CO COOH + N14H2

Thus, although this type of labeling permits following the course of the nitrogen metabolism, it is valueless as a guide to what bappens to the remainder of the molecule after deamination; for this purpose, the carbon chain should contain isotopic earbon, preferably in as many positions as possible, or stably bound deuterium may be present.

Measurements of blood volume and extracellular fluid volume have long been made by nonisotopic dilution methods involving the injection of some substance foreign but harmless to the body. A known amount was injected, time allowed for equilibration, and the concentration measured in the blood plasma. With a substance such as the dyestuff T 1824 (Evans blue), which escapes but slowly from the blood vascular system, a close estimate of blood plasma volume is possible. With soluble bromides or thiocyanates, which diffuse through all extracellular fluids but do not significantly enter cells, extracellular fluid volume can be measured. For this latter type of measurement the use of radioactive Br permits the use of very small doses and avoids the pharmacological effects of this ion. For blood volume determinations, the use of red cells tagged with radioactive phosphorus obviates the errors inherent in the dye method For measurement of total body water, deuterium oxide is the indicator of first choice, since it mixes freely with hody water and traverses all cell houndaries. Its chief disadvantage is the special instrumentation required for its measurement (see "Stable Isotopes," below) Heavy water is toxic, but not in the low concentrations (0 2 atoms per cent deuterium) used in such studies A minor error arises from exchange of deuterium and from its incorporation into organic molecules. This error has been estimated at less than 1 5 per cent. 10

EXPERIMENTS ON RADIOACTIVE ISOTOPES

Determination of Blood Volume: Principle. The determination of blood volume in rabbits as accomplished by neubating blood cells from the animal with 80 to 100 microcuries of P²¹ labeled phosphate for 45 to 60 minutes at 37° C. Approximately 10 per cent of this activity is absorbed by the cells, which are then washed, reconstituted to volume, and impeted into the animal. After an interval for distribution of the labeled cells throughout the circulation, a blood sample is withdrawn for counting. The dilution of the activity of the necessary of the volume of the whole blood is obtained by dividing the activity of the injected cells by that of the blood wildrawn.

The relative simplicity of the experiment allows for total working space on one 24-inch by 30-inch table. This is covered with a large blotter so that any spilled active material may be readily discarded. Special apparatus consists of a basic scaler and Geiger-Muller counter with a shielded manual sample changer. The counting apparatus is kept on a table separate from the working area. The radiation hazard is

^{**} Edelman Olney James Brooks and Moore Science 115, 447 (1952).
** Kelly, Simonsen, and Elman J. Clin. Invest., 27, 795 (1948), modified for use with rabbits by Dr. Paban Lionetti.

minimal to the animal or to the experimenters, as shown by Reid and Orr, 12 who give data for 100 humans with average uptake of the cells amounting to 7 per cent of the 50 to 100 microcuries incubated, with the largest single uptake 19 5 per cent. In calculating the bazard when the activity was injected into a burnan, they assumed a 20 per cent uptake in a 50-kg individual. This would calculate to 0.07 rep for approximately one week, which would diminish to 0 035 rep in two weeks (14 3 days balf life) This calculated value is deliberately high, for it assumes no excretion, but is yet a small fraction of the normally permissible 0 3 rem With but small precaution. therefore, no radiation bazard is eccountered Indeed, it is probable that the students are exposed to greater dangers in their other experiments by handling corrosive acids and hot solutions. As a means of illustrating fully the manner of working with isotopes students may characterize the radiophosphorus they use by calibrating its millicurie strength against a set of simulated reference standards. They may also count a sample daily for two minutes for seven or more successive days, in order to calculate the half his of the P12 used and thus ideotify it This also affords a check on the purity of the isotone shipment, and acquaints the student with the main remaiolog features of low-energy isotope experimentation

Procedure Ail glassware should be siliconed and dried To all Cutler or ,imilar tubes, a pinch of heparin (gauge by tip of spatula) should be added

- (a) PREPARATION OF TAGGED CELLS To two 15-ml centrifuge tubes approximately 1 5 to 2 mi of acid citrate destrose solution (ACD)12 is added. Using a syringe wet with ACD, 20 to 30 ml of blood is taken from a rabbit by cardiac nuncture Between 5 and 8 ml of the blood is placed in one and the rest in the other centrifuge tube. These are covered with parafilm and inverted 20 to 30 times to prevent clotting Both tubes are placed in the incubator at 37° C. until ready for use, and inverted Intermittently To the tube containing 5 to 8 mi blood, add 80 to 100 microcurles Parla volume less than 1 mi (to maintain Isotonicity) The tubes are inverted 5 to 10 times and returned to the incubator for 45 to 60 minutes. Invert the tubes 10 times approximately every 5 minutes Care is required to avoid shaking the tubes At the end of the incuhatlon time, the cells are spun down in a refrigerated centrifuge at 2000 r p m for 15 minutes The fluid is removed with a capillary pipet and replaced with cold ACD to the original volume Recentrifuging and reconstituting to volume is repeated 3 times, mixing belne done by careful inversion 20 to 25 times If slight hemolysis is observed during the washing, it is advisable to eliminate the final one or two washings After removal of the third washing, the volume is replaced to original with cold ACD. The tube is inverted gently 20 to 25 times or more, and placed in the icebox until ready for use
 - (b) PREFARATION OF THE TAGGED CELLS FOR COUNTING INITIAL ACTIVITY Since the tagged cells are too active to count conveniently, they are diluted serially with inert blood. Into each of 4 small test tubes containing heparin place 3 mi of blood. Care is required to prevent clotting. The pipet should be previously rinsed with ACD. To the first tube is carefully added exactly 1 ml of the tagged red cells (making sure that the cells are well mixed before the addition). The tagged cells in the first tube have thus been diluted 4 times

One mi of the 1-4 suspension in the first tube is now removed and added to the second tube containing 3 ml of untagged blood, this gives a 1-16 dilution Transfers are continued until the fourth tube is prepared containing a 1-256 dilution of the initial 1-ml of tagged cells. Mix carefully by drawing blood into and expelling it from the pipet 15 to 20 times. A pair of 1-ml portions of the final dilution is placed in the planchet for counting. The each is added a drop of 1 per cent detergent solution (Duponol) to facilitate spreading in the planchet.

(c) INECTION OF RADIOACTIVE CLUS. The rabblt is weighed and placed in a box with a cover so notched as to allow the head and ears to protrude Both ears are shaved along the marginal ear veins. The tagged cells are removed from the leebox and inverted to mix them well One mi is drawn into a tuberculin syringe and all air bubbles are removed. Holding the right ear tightly, introduce the needle into the right marginal veln. Once in the veln, the needle is extended to the hilt. If the needle sildes easily with little fric tion, the left thumb is placed over the needle and held securely, in order not to withdraw the needle if the animal moves its head Inject the cells slowly They are infused readily if the needle is in the vein The needle is then with drawn and the site clamped with the hulldog clamp After 20 minutes of equilibration in the rabbit, a sample is obtained for counting hy making a nick in the left marginal ear vein Preliminary treatment with xylol or toluene on the surface of the ear may be used to promote a more rapid flow Several mi of blood are collected in a heparinized tube and 1 ml is pipetted into a planchet Five drops of Duponol (ca 1 per cent) is added The planchet of diluted tagged cells (1 256) and the one prepared from the animal are incubated at 37° C to dryness (overnight) A third sample of untagged whole blood is similarly prepared for taking background counts If it is necessary to save time, the wet samples may be counted, but the counts are less repro ducible and less accurate

(d) PACKED CELL VOLUME This part of the procedure is necessary only if plasma volume is to be determined. The plasma volume is found from the whole blood volume by a hematocrit determination, finding the plasma fraction of the whole blood volume. The tube with the remainder of the blood is spun at 2500 to 3000 r p m for 60 minutes. The top level of the plasma at the meniscus is read. The level of the packed red cells is noted, ignoring the buffy coat. The second declinal is estimated in the read with the packed red coat.

CALCULATION The instrument background B is measured in counts per minute using the sample of mert blood. The background count B is subtracted from the counts of both the experimental and injected samples. The experimental samples withdrawn from the animal are counted for a time interval corresponding to secrat thousand counts and the activity in counts per minute is recorded (E) after subtracting B. The activity of the injected sample I is found from the count (corrected for background) of the diluted tagged blood D multiplied by the dilution factor 256 thus

$$I = D \times 256$$

The wlole blood volume (WBV) in in likiters is therefore

 $WBV = \frac{I}{E}$

RADIOACTIVE ISOTOPES IN MEDICINE

Ionizing radiation has been applied with increasing skill and success to destroy tumors or other undesired tissues. Radioactive isotopes in

general are applicable as sources of ionizing radiation for this purpose. Certain radioactive isotopes are concentrated by physiological mechanisms in certain cells or tissues—I¹³¹ in the thyroid, P³² in blood cells—just as the normal stable isotopes of these elements are so concentrated. In this manner, ionizing radiation can be localized to certain tissues for their destruction when they have hecome neoplastic. After administration of P³² by mouth or intravenously (in the form of a phosphate), it is taken up first by the red cells, with concentrations in white cells increasing after the second day. In leukemic patients who have received P³², after one week the isotope concentration in lenkocytes is four to five times that in red cells.¹⁴

Smaller doses of radioactive isotopes may he used in diagnostic studies. The amount of 1³¹ in the thyroid gland of a human patient can be estimated by a measurement made externally over the thyroid region by a 4-tuhe Geiger-Muller counter. In If an oral tracer dose containing 100 to 150 microcuries of 1³¹ is given, 30 or 40 per eent of it should be demonstrable in the thyroid gland ou the next day in patients with normal thyroids. Hyperthyroidism is characterized by higher uptake of iodine—around 70 per cent, while in hypothyroidism the uptake is low—around 12 per eent as measured under similar conditions. Administration of 1¹³¹ in larger amounts is utilized for the destruction by radiation of cancerous thyroid tissue, or in reducing the activity of the gland in hyperthyroid patients. This form of treatment hrings ahout results comparable to surgical removal of the thyroid, in whole or in part according to the dosage of the radioactive material.

STABLE ISOTOPES

Most of the common elements as ordinarily encountered in nature consist of mixtures of stable isotopes. Thus ordinary oxygen consists of the three stable isotopes O16, O17, and O18, in the proportion respectively of 99.76, 0.04, and 0 20 per cent. More than 200 stable isotopes of the various elements have been recognized as existing, but only a very few have been obtained in concentrated form. To obtain the separate isotopes from a mixture, or fractions relatively enriched with respect to one isotope, advantage is taken of properties such as diffusion or reaction velocity which may vary with mass differences. Thus a water molecule containing O18 will be heavier than one containing O16, and the two types may be separated by controlled fractional distillation. By such methods or their conivalent, the stable isotopes H2 (deuterium), C12, N15, O15, and S14 have been made available for biological investigation. The first of these to become available in amount sufficient for investigative purposes was deuterium, first obtained by Washburn and Urey m 1932 by the fractional electrolysis of water. It was in fact the availability of deuterium which suggested to the imaginative genius of Schoenheimer and Rittenberg the possibility of its use in biological investigations, a concept which may be said to have initiated the present phase of application of isotopes to biological problems.

Duffy and Howland N. Y. State J. Med., 52, 551 (1952).
 Freedberg, Chamovitz, and Kurland. Vetabolism, 1, 20 (1952).

The cluef application of the stable isotopes in biological investigation is for the purpose of labeling or tagging a molecule or part of a molecule. The stable isotopes have the advantage over the radioactive type in that the time of preparation of the compound incorporating the isotope, or the duration of the experiment, are of no importance, there is no question concerning the possible effect of radiation on the experiment, and in some instances the isotope may be recovered at the end of the experiment and used again. Disadvantages relative to radioactive isotopes include the increased difficulty of measurement, the necessity for a higher concentration, since measurements of radioactivity are far more sensitive than measurements of mass, and the relatively few stable isotopes available.

With the exception of deuterium, the stable isotopes are measured with an instrument ealled the mass spectrometer. The principle upon which this instrument operates is illustrated diagramatically in Fig 259 The sample in the form of a gas, e.g. N2, CO2, is admitted to a highly evacuated chamber across which a stream of electrons is flowing An electron colliding with a gas molecule causes it to become ionized, e.g. CO2 → CO2 The gas ions so formed are accelerated by suitable means to produce a stream of ions which passes along the tube shown and through a powerful magnetic field, so arranged as to deflect the ion from its initial path On leaving the magnetic field, the ions pass to a collector plate and produce an ion current which is what is actually measured as an index of the number of ions striking the plate Ions of different mass, eg C12Of and C12Of, may be differentiated by either varying the strength of the magnetic field, or more commonly this strength is held constant and the accelerating voltage is varied Results are obtained in the form of a graph relating ion current to accelerating voltage Isotopie ions differ on such a graph in their position along the voltage scale (which can be converted into mass, since a constant relation exists between these two quantities), the ratio of one isotope to another is given by the ratio of ion currents at the respective points on the mass (voltage) scale It is this ratio of isotopes which is ordinarily desired-i e, the relative abundance for example of C12 over C13 in the sample as compared to their relative amounts in ordinary nonisotopie compounds

The measurement of deuternum is a special case It may of course be measured with the mass spectrometer, but equally satisfactory results can be obtained by converting the deuternum to the form of water (as by burning the sample in oxygen) and determining the specific gravity of the water so produced. There is a predictable relationship between the specific gravity of the water and its content of heavy water, D₂O A common method for determining specific gravity is the falling drop method, in which the time required for a drop of water to fall through a definite distance in a nonaqueous solvent is measured. This method is capable of light accuracy.

nigh accurac

The isotope content of compounds labeled with the stable isotopes usually is expressed in terms of atoms per cent of isotope. For example, if a substance contains 12 hydrogen atoms, and one of these is replaced by deutenum the isotopic content will be 1/12, or 8.3 atoms per cent

This is subject to some correction, since an ordinary "nonisotopic" compound contains a certain proportion of isotopes in accordance with their normal abundance. The excess of a given isotope over that normally present is referred to as aloms per cent excess, a more subjectory term when dealing with variations in the isotope content of compounds

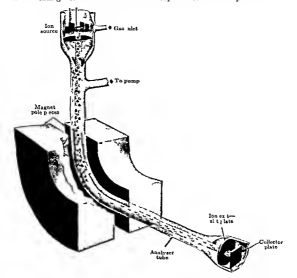


FIG 259 OPERATION OF THE MASS SPECTROMETER TUBE IDEALIZED VIEW.

The ion beam is projected into the tube and deflected by the magnetic field.

The separation of the ions of different masses is indicated by the variance in the radius of curvature of the gray and black arrow beams in the area of the magnetic field.

Courters Westinghouse Engineer

A special problem arises in connection with the use of deuterium in metabolic tracer studies. Deuterium has twice the atomic weight of ordinry hydrogen and hence diffuses more slowly and will slow down reaction rates. Differences in atomic weight cost (by definition) among all isotopes of a given element but in no other instance is the difference nearly as great proportionately.

Analysis by Isotope Dilution Stable or radio active isotopes may be used in chemical analysis in just mees where other methods fail or

account of lack of specificity, as for example in the assay of culture filtrates of Penicillium chrysogenum for penicillin G to the exclusion of other penicillins. Four steps are involved in such an isotope dilution assay: (a) a tracer sample of the compound to be determined (e g., penicillin G) is prepared with a known isotopic content, and with the isotopic atom in a position which will not exchange; (b) a known quantity of this isotopically tagged compound is isolated in a state of purity, but not necessarily quantitatively; (c) the isotopie content of the isolated sample is determined. By comparison of this figure with the isotopic content of the tracer sample prepared in step (a), and knowing the proportions in which the mixture in step (b) was made, the quantity of the substance in the unknown sample (eg, penicillin G in the culture filtrate) can be immediately calculated. If the isolated sample has the same isotopic content as the tracer sample, the quantity in the unknown was zero If the isolated sample has half the isotopic content as the tracer, the quantity in the unknown was equal to that in the tracer added; if onethird, equal to twice that added; one-fourth, equal to three times that added, and so on. In general, if the isotopic content of the isolated sample is x per cent of the original, the quantity in the unknown is of the quantity added.

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1951

Carbohydrate, Fat, and Protein Metabolism

GENERAL

The components of a normal mixed diet which supply all the requirements of the animal body for growth and maintenance include proteins, fats, carbohydrates, the various vitamins, water, and certain morganic elements. In the present chapter are considered some of the nutritional requirements for proteins, fats, and carbohydrates and the changes which these substances undergo subsequent to their digestion and absorption into the animal hody. The energy metaholism of these substances has heen considered in Chapter 25. Brief discussions concerning certain metaholic phases are also found in the chapters dealing with the urmary constituents and in other connections. The questions of mineral metabolism, water, and the vitamins are discussed in subsequent chapters.

Throughout the discussions which follow, frequent mention will be made of the use of isotopes, which have proved such valuable tools in the study of intermediary metabolism and nutrition. The isotopes themselves and their use in hology are discussed in Chapter 32. Adequate entrance to the voluminous literature of intermediary metaholism and nutrition sufforded by the footnote references and the reviews and special articles.

listed in the bibliography at the end of this chapter

CARBOHYDRATE METABOLISM

Carhohydrates supply the major portion of the daily energy requirements of the normal individual, on an ordinary diet more than half of the total daily calories usually come from this source. In addition to being oudized as a source of energy, carbohydrates may be transformed to glycogen, supply the carbon chain for certain amino acids, or be converted into fat. Of these various processes, glycogen formation and breakdown ap-

pears to occupy a central position

Glycogen Formation and Breakdown (Glycogenesis and Glycogenesiyss). Glycogen is found in practically all of the tissues of the body, but in varying amounts, the glycogen content of the brain, for example, is so low as to require special care for its detection. The muscles and liver contain most of the glycogen of the body, largely because of their bulk relative to thirt of the other tissues. Glycogus from various tissues—e.g., muscle glycogen, hive glycogen—appear to be chemically similar if not identical but to have markedly different physiological significance. Thus the glycogen content of muscle is relatively constant in the about a

of exhaustive inuscular contraction and is little affected by starvation of the nature of the diet, which may produce variation in the liver glycogen content from a mere trace to upward of S per cent of the weight of this organ

Glycogen, therefore, may have both a functional and a storage significance. The capacity of the animal to store carbohy drate as glycogen is relatively limited, however, as contrasted with its capacity to store fat. The adult human body may contain about 300 g of glycogen, of which only that fraction found in the liver (normally about one-half of the total) represents a significant source of carbohy drate for general metabolic purposes. Fasting for a few days is usually sufficient to deplete the aumal of all available stored carbohy drate. Any carbohydrate metabolism in a fasting animal after the exhaustion of stored carbohydrate is due presumably to synthesis of carbohydrate from nonearbohydrate precursors (gluconeogenesis)

Major precursors of liver glycogen within the animal body include the glucose fructose, and galactose (but not pentose) produced by carbohy drate digestion in the intestinal tract, the glucose and lactate of the blood the glycerol portion of the fats of either the diet or the body, and certain amino acid constituents of the proteins of either the diet or the tissues which are capable of heing converted into glucose and hence into glycogen Precursors of muscle glycogen include the glucose of the blood and, to an extent which is still debatable, the lactic acid produced from glycogen itself during muscle contraction. These various interrelationships with regard to liver and muscle glycogen may be summarized in the following diagram adapted from Con.

Carbohy drate (det)
Glyccrof from fat (det and body)
Glycogenc amino acids (det and body)
Glycogenc amino acids (det and body)
Blood lactic (lact and body)
acid acid acid

Knowledge that a particular substance is a precursor of liver gly cogen, for example usually has been obtained by demonstrating that an in creased liver glycogen content follows the feeding of the test substance Perfusion of the isolated organ and incubation with surviving liver slices also have been used It has generally been assumed that if glycogen for mation occurred, the test substance was directly incorporated into the newly formed glycogen That the situation is not quite as simple as this has been unequivocally demonstrated by the use of glycogen precursors labeled with radioactive carbon For example, if lactate labeled with radioactive carbon is fed to animals whose liver glycogen has been depleted by fasting and the newly formed glycogen which results from the lactate feeding is subsequently isolated only a fraction of this glycogen proves to contain the isotope label Furthermore the isotope is found distributed among the various carbon atoms of the glucose residues in the labeled gly cogen in such a way as to indicate that the fed lactate had entered into other reactions before some of the carbon atoms became incorporated into glycogen. Similar results have been obtained with glucogenic amino acids labeled with isotopic carbon. If glycogen formation is stimulated by glucose or lactate feeding in the presence of radioactive carbon dioxide (as hicarbonate), the radioactive carbon is also found in the glycogen suhsequently isolated. Thus glycogen formation and breakdown is a complicated and continuous process within the hody; according to Stetten, 't the half-life of rat-hver glycogen is but one day. The accumulation of glycogen in the presence of an apparent precursor may he due only in part to direct utilization of the precursor itself, or may even represent a "sparing" action on glycogen metabolism without any direct connection with such metabolism.

Knowledge concerning the intermediate steps in glycogeu formatiou and hreakdown has been obtained largely with tissue preparations or isolated enzyme systems. Much of this work has heen done with either muscle or liver; it is helieved that conclusions drawn from such studies are generally applicable, in principle if not in specific detail. The various reactions known to be concerned in glycogen formation and breakdown may be divided into two phases, auaerobic and aerobic, summarized as follows:

Glycogen + Phosphate

Glucose-1-phosphate

Glucose + ATP → Glucose-6-phosphate → Glucose + Phosphate

Fructose + ATP → Fructose-6-phosphate

Il

Fructose + ATP → Fructose-1,6-diphosphate

Il

Triose phosphates

Il

Phosphoglycerate

Il

Phosphopyruvate

Many of these reactions already have been discussed in detail in Chapter 10 in connection with the chemistry of lactic acid formation in muscle, and the chemical structures of the various intermediate compounds also will be found there.

Lactate Pyruvate Anaerobie phase

1 ----
CO₂ + H₂O Aerobic phase

It will be noted that the immediate ehemical precursor of glycogen is the compound glucose-1-phosphate (the Cori ester). The enzyme catalyzing the reversible reaction between glycogen, inorganic phosphate, and glucose-1-phosphate is known as phosphorylase. The equilibrium in this reaction is in favor of glycogen; the direction of reaction appears to be determined largely by the concentration of inorganic phosphate. In the presence of excess inorganic phosphate, glycogen breakdown occurs; for

¹ Stetten and Boxer: J. Biol. Chem., 155, 231, 237 (1944).

glycogen synthesis, it is necessary to keep the inorganic phosphate contentiow. This apparently is done in the cell by oxidative processes which incorporate the phosphate into phosphate esters, some of which are intermediates in the scries of reactions shown. There is thus in effect a phosphate cycle (Cori), whereby inorganic phosphate liberated by glycogen formation is used to phosphorylate other compounds which, in turn, can

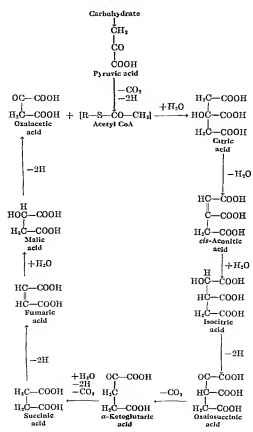
give rise to glycogen and morganic phosphate again. For free glucose (or fructose) to be converted into glycogen by the reactions shown, phosphorylation is necessary. This phosphorylation—leading to the formation of either glucose-6-phosphate or fructose-6-phosphate, as the case may be—requires the presence of the "high energy phosphate" of adenosinetriphosphate (ATP), and of enzymes of the recokinase, similar enzymes are known as glucokinase and fructokinase, similar enzymes are found in muscle. In hrain tissue there appears to be only one enzyme which, like yeast hexokinase, acts upon both glucose and fructose. The hexokinase enzymes, or at any rate their activities, are apparently under the control of certain hormones, as discussed on p. 1000, and knowledge concerning this control may prove to be of fundamental importance in understanding certain phases of carbo-

livdrate metabolism

After phosphorylation, glueose becomes available for either glycogen formation, reconversion to glueose, or hreakdown to lactic acid anaero-incally or complete oxidation aerobically Reconversion to glueose is due to the enzyme phosphatase, which catalyzes hydrolysis of glueose-6-phosphate to glueose and inorganie phosphate. This is believed to be the origin of the glueose of the blood from the glycogen of the hiver (and possibly also of the kidneys). In muscle tissue, which contains no phosphatase, glueose-6-phosphate is either converted to glycogen or degraded to form lactic acid anaerobically or completely oxidized aerobically. The formation of lactic acid from glycogen requires no oxygen, the oxidation of trosephosphate to phosphoglycerate is coupled with the simultaneous reduction of pyruvate to lactate through the mediation of conzyme is which is alternately reduced and oxidized as described in Chapters 10 and 35.

Aerobic Oxidation of Carbohydrate. The precise mechanism whereby carbohydrate is completely oxidized to CO₂ and H₂O is still a subject of discussion it is felt by many that a reasonable answer to the mechanism of acrobic carbohydrate oxidation is afforded by the cycle process first discribed in its essential principles by Krebs and called by limith catter acad cycle later modified by Krebs and others and now more

generally known as the trearboxylie acid cycle
According to this concept carbohydrate is considered to be degraded
to the stage of pyruve acid CHLOCOOH by the series of reactions
just described for the aucrobic breakdown but whereas under anacrobic
conditions pyruvic acid is concerted to factic acid as the chief end product
of the process under acrobic conditions the pyruve acid undergoes a
different series of reactions. These reactions of the tricarboxylic acid cycle
art, summarized in the accompanying diagram which for purposes of
clarity has been considerably simplified as will be pointed out.



As shown in this diagram the first major reaction in the acrobic oxidation of pyrruvic acid is the production of citize acid from pyrruvic acid and oxalacetic acid Earlier doubts concurring the role of citize acid tiself in the citize acid cycle have now heen resolved and there is general agreement as to its importance as indicated. Oxalacetic acid is normally present in small amount in actively respiring tissue and is constantly being recentated as will be shown.

The exact mechanism of the formation of citric acid from pyruvic and oxalacetic acids has been the subject of intensive study Earlier views based on a variety of evidence postulated the presence of a reactive 2 carbon fragment (active acetate) formed from pyruvic acid which condensed with oxalacetic acid to form citric acid. Active acetate has now been identified by Lymen as the S acctyl ester of coenzyme A acetyl CoA symbolized as follows R—S—CO—CII, where R is the remainder of the coenzyme A molecule (see pp. 1190-1

The mechanism of formation of acetyl Co\ from pyruvic acid in animal tissues is not entirely clear in all its phases but there is no doubt that it occurs in fact it is believed that all substances entering the citne acid cycle at the 2 carbon fragment level must be precursors of acetyl Co\ One method of formation of acetyl Co\ from pyruvic acid which has been established for heart tissue by Korkes et al is the reaction between pyruvic acid coenzyme \ and diphosphopyridize nucleotide (DP\), coenzyme I) in the presence of diphosphothiamine (cocarhoxylase), to give acetyl Co\ Co\ 2 and reduced DPN

Pyruvic acid + DPN + CoA \rightarrow Acetyl Co \ + CO₂ + DPNH + H⁺

In the presence of the condensing enzyme which is the first enzyme of the citric acid cycle to be isolated from animal tissues in crystalline form (Ochoa) acetyl Co's readily condenses with oxalacetic acid to form either acid and Co's

Acetyl Co 1 + Oxalacetic acid → Citric acid + CoA

Thus the over all reaction results in the formation of citric acid from pyruvic acid (by way of acctyl CoA) and oxalacetic acid The role of DPN is that of hydrogen acceptor in the living cell the hydrogen is ultimately transferred to oxygen. The role of occarboxylase is not entirely clear A similar oxidation of pyruvic acid can be demonstrated in certain bacterial extracts here the occarboxylase is said to be associated with 6.8-thioctic acid (see Chapter 33) in its action

Citrie acid is readily metabolized in muscle and other animal tissues through the stages of cis aconitic acid and isocitrie acid as shown in the presence of the enzyme aconitase which mediates an equilibrum among the three compounds Isocitrie acid once formed undergoes oxidation by dehydrogenation in the presence of isocitrie deliydrogenase to yield oxidosuccinic acid which is then decarboxylated to give a ketoglitarie acid It is not clear whither an oxidosuccinic decarboxylase is concerned here or the entire reaction from isocitrie acid to a ketoglitarie acid is

² Korkes del Camp llo and Ochoa J Biol Chem 195, 541 (1952)

mediated by isocitric debydrogenase alone It is known however that triphosphopyridine nucleotide (TPN, coenzyme II) is coupled in these reactions, so that the oudation of isocitric and involves the reduction of TPN. The over-all reaction at this particular step is therefore as follows.

Isocrtric acid + TPN $\rightleftharpoons \alpha$ -Ketoglutane acid + CO₂ + TPNH + H+

As in the previous reaction involving DPN, the hydrogen of the reduced

TPN is ultimately transferred to oxygen to form water

The substance α -ketoglutane and is readily converted by decarbo-ylation and dehydrogenation (with DPN as hydrogen acceptor) to give the C4 dicarbo-ylae and succeime and The intermediate stages in this reaction are not entirely clear, but they appear to involve the formation of succinyl CoA as an intermediate. Succinic and is then acted upon by the enzyme succeime dehydrogenase, present in most animal tissues, to give fumance and by oxidative dehydrogenation. Fumance and readily undergoes bydration in the presence of the enzyme fumance to form malic and, which on oxidation in the presence of the enzyme make dehydrogenase yields oxidacetic and At this point the entrance of another molecule of nyruve acid permits the entire evelot to be repeated

Thus the not result of the cycle is the complete disappearance of one molecule of pyruve acid, giving rise to three molecules of CO₂ in the process and requiring the ultimate presence of five oxygen atoms (25O₂) to accept the 10 hydrogen atoms removed by the various dehydrogenation reactions. As indicated above, the mechanism of bydrogen transport to oxygen is by way of the di- and triphosphopyridine nucleotides (coenzymes I and II), the flavoproteins, the cytochromes, and cytochrome

oxidase, as described in Chapter 12, "Enzymes"

The over-all reaction is therefore

 $CH_4 CO COOH + 25O_2 = 3CO_2 + 2H_2O$ Pyruvic
acid

with a respiratory quotient (R Q) of 3/2.5 = 1.2 By the combination of reactions described, involving the formation of pyruvate from earbohydrate and the participation of pyruvate in the triearboxylic acid cycle, it is possible therefore to account for the complete oxidation of carbohydrate to CO₂ and H₂O. To obtain an R Q of 1.0, which is the R Q of earbohydrate oxidation, the transfer of two more hydrogen atoms to an oxygen atom must enter into the picture. These two hydrogen atoms are presumably those which arise in the oxidation of triosephosphate during the anaerobic phase of earbohydrate breakdown (see p. 274), and which produce lactic acid from pyruvic acid under these conditions. In aerobic metabolism they are transferred to oxygen by way of diphosphopy ridine nucleotide. (DPN)

The major evidence in favor of such a cyclic process as that described may be summarized as follows the various postulated intermediates usually increase the respiratory rate of sintable tissue preparations (c.g., mineed muscle) when added in citaly the amounts, many of the individual

enzymes concerned bave been isolated and shown to function as indicated certain of the postulated intermediates have been isolated from tissue systems in the presence of specific poisons such as malonate and arisente and critical examination of the theory by the use of isotopes as markers for specific portions of the various molecules concerned as yet has not revealed any major discrepancies

The reactions of the tricarhoxylic acid cycle afford an explanation for the participation in the processes of earlichydrate exidation of the various intermediates shown even though such intermediates are not earlichydrates or may arise from nonearhohydrate precursors. The implications of this relationship to the metaholism of fats and amino acids will be evident from the subsequent discussion in this chapter.

Most of the reactions of the tricarboxylic acid cycle are reversible in explanation is afforded for the synthesis of carbohydrates from non carbohydrate precursors which are or giverse to components of the cycle provided a means is available for their conversion to pyruvate. There is ample evidence that such formation of pyruvate occurs in amma tissues Pyruvic acid may arise by the direct decarboxylation of oxalaceth acid.

COOH CH2 CO COOH - CH2 CO COOH + CO2
Oxalacetic acid Pyruvic acid

This reaction is known to occur in animal tissues. Once formed pyruvi acid is reachly pho phorylated in the presence of ATP

Pyruvate + ATP

⇒ Phosphopyruvate + ADP

Phosphopyruvate then can serve as a direct precursor of the hexoses an glycogen hy the reversal of the glycolytic reactions already described c p 989 Kalckar has shown that phosphopyruvate itself is formed durf the aerobic oxidation of the C4 dearhoxytic acids hy tissue extracts the may arise by the decarboxylation of phosphooxalacetic acid (Lipmann,

COOH CH CO(PO₁H₂) COOH → CH₂ CO(PO₂H₂) COOH + CO₂

Phosphooxalacetic acid Phosphopyruvic acid (enol form) (enol form)

(enol form)

Other reactions doubtless remain to be discovered but at any rate it be clear that the pathways of carbobydrate synthesis from noncarboby drate precursors are open by the reactions described

The possible reversibility of the decarboxylation reactions in the citric acid cycle has stimulated interest in the mechanisms of CO, fixation 16 the utilization of the carbon of carbon dioxide for metabolic purposes. That CO, fixation occurs elsewhere than in green plants and photosynthetic bacteria was first shown for nonphotosynthetic bacteria by Wood and Werkman who postulated that the reaction molyled the formation of oxalacctic acid from pyruvic acid and carbon dioxide

CH₂ CO COOH + CO₂ → COOH CH₂ CO COOH Pyruvic acid Oxalacetic acid

Proof that CO2 fixation occurs in animal tissues was afforded by Evans

and Slotin² using bicarbonate containing radioactive carbon, the \(\alpha \)-keto glutaric acid synthesized by pigeon-liver preparations in the presence of pyruvate and the isotopic bicarbonate was found to contain a significant amount of the isotopic carbon. The carbon of radioactive sodium bicarbonate administered to rats is also found in the glycogen subsequently isolated from the liver ⁴. When fatty acids are oxidized to acctoacetate by liver preparations in the presence of radioactive carbon dioxide, the isotope is found incorporated, to a small but significant extent, in the carboxyl group of the acctoacetate formed ⁵ Other examples could be cited, they all demonstrate that the release of carbon dioxide by animal tissues is not an evclusively one-way process.

Although the Wood-Werkman reaction has been much studied in connection with CO₂ fixation by animal tissues, there is no good evidence that it occurs elsewhere than in bacteria Of greater interest in this connection is the reaction discovered by Ochoa in liver extracts, in which make acid is reversibly outlized and decarhoxylated to give pyruvic acid and CO₂. This reaction requires TPN as hydrogen acceptor, and is

catalyzed by the malic enzyme

Malic acid + TPN

⇒ Pyruvic acid + CO₂ + TPNH + H+

If this system is coupled with another TPN-requiring system, such as the combination of glucose-6-phosphate and glucose-6-phosphate dehydrogenase, the TPN may be removed from the system to such an extent that the reaction proceeds to the left and there is a net synthesis

of malic acid from pyruvic acid and carbon dioxide

It is of interest to note bere that Ocboa had previously shown that the oxidative decarboxylation of isocitine and to form a ketogliutant acid was experimentally reversible under the proper conditions, bowever, the equilibrium conditions for the make enzyme are stated to be 15 times more favorable for CO₂ fixation by the make enzyme reaction than for the isocitrate-ketogliutarate reaction. It is to be noted that, once make and is formed from pyruvie and and earbon dioude, the presence of make dehydrogenase in a tissue will lead to the production of oxalacetic acid. Thus the make enzyme provides a mechanism for the entry of pyruvie and into the entre acid eyele other than by way of acctyl CoA

Despite the undoubted demonstration of its existence, the significance of CO₂ fivation by aumal tissues remains obscure. The extent of fivation is usually quite small, since equilibrium conditions ordinarily favor decarboxylation. The comparative ability of various animal tissues to fix CO₂ has been studied by Crane and Ball * According to Ochoa, the malic enzyme reaction provides the only means thus far known whereby a net synthesis of dicarboxyle needs can be obtained from pyruvic acid and carbon droxide. It has been suggested that CO₂ fixation provides for the constant presence of the eatlythe amounts of dicarboxyle acids neces-

Evans and Slotin J Biol. Chem. 136, 301 (1940)
 Solomon Vennesland Klemperer Buchanan and Hastings J Biol Chem. 140, 171, 341)

¹ Haut and Lardy J Bud Chem 192 435 (1951)
Crane and Ball J Bud Chem 188 S19 (1951)

sary for the continuous operation of the eitric acid cycle. It is interesting to recall here that, as discussed in Chapter 2, the assimilation of CO₂ by green plants during photosynthesis leads to the formation of pho phoglyceric and phosphopyruvic acids, and that malic acid is a subsequent product of the reaction. Thus the fundamental processes of CO₂ assimilation in plant and animal tissues may well be similar.

Despite the extensive knowledge which has been obtained concerning intermediary carbohydrate metabolism, there is much that is still obscure The aerobic oxidation of lactic acid, for example, usually is con sidered to occur by way of conversion to pyruvic acid which is then oxidized by the processes just described. There is some evidence that other as yet unknown and possibly important pathways exist. In certain tissues, glucose can be shown to be oxidizable to form glucome or phosphoglucome acid, and the latter has been shown to be converted into ribose phosphate, one of the building blocks of nucleic acids. The conversion of carbohydrate to pyruvate and the reactions of the tricarboxylic acid cycle have been studied largely in one tissue, i e, muscle, and the con clusions derived are not necessarily applicable in toto to other tissues Yet Krebs has pointed out that the various animal tissues thus far investi gated possess enzymes similar in general to those required for the tn carboxylic acid cycle, so that while individual differences between tissue doubtless occur, the basic reactions could be fundamentally similar in al tissues

Formation of Carbohydrate from Ammo Acids The reactions of the thearboxyhe acid cycle afford a reasonable explanation for the ability of certain ammo acids to give rise to glucose or extra glycogen in the ammo acids to the second acid appartic acid, and alanine, by metabolic deamination, give rise to α-ketoglutane acid, and acetic acid, and privive acid respectively. These three latter compound are recognized components of the thearboxyhe acid cycle

glutamic acid → α ketoglutame acid aspartic acid → oxalacetic acid alamine → pyruvic acid

After deamination therefore these three amino acid constituents of the protein molecule become indistinguishable from carbohydrate metabolites. Since the reactions of the trachoxyle acid cycle are reversible, a mechanism is available for either the synthesis of carbohydrate or metabolism via carbohydrate pathways for these amino acids, and for any others which are convertible into these amino acids by metabolic processes. That pathways exist other than those described is quite probable, but they remain to be discovered.

Conversion of Carhohydrate to Fat. The fattening of farm animals and the development of obesity in man on high carbohydrate diets suggest that carbohydrate can be converted to fat in the animal body in feeding experiments in which a control animal is killed and analyzed for fat while a litter mate is fed a high carbohydrate diet, it can be shown that the latter may after a time contain more fat than could be derived from all of the fat and protein fed. In the same way, milch cows without

loss of fat from their bodies may secrete in their milk much more fat than can be accounted for by fat and protein ingested. Studies of the respiratory quotient yield similar evidence with regard to the couversion of carbohydrate into fat (see Chapter 24). The availability of isotope-labeled glucose has provided a more direct demonstration of the conversion of carbohydrate to fat. For example, if mice and rats kept on an essentially fat-free diet are given a dose of labeled glucose, between 10 and 15 per cent of the labeled carbon atoms are found in the fatty acids of the animal body after 24 bours. These and other results lead to the conclusion that conversion of carbohydrate to fat does not represent merely the storage of energy, but is also an important pathway in the metabolism of carbohydrate itself.

The glycerol for fat synthesis may arise very easily from carbohydrate, possibly directly from phosphoglyceraldehyde, a normal intermediate in earbobydrate metabolism in tissues:

CHO H³COH H³COH	
HCOH — HCOH — HCOH → HCOH	
H ₂ COPO ₂ H ₂ H ₂ COPO ₂ H ₂ H ₂ COH -	⊥ н.РО.
Phosphogly- Phospho- Glycerol ceraldehyde glycerol	11111 04

The mode of synthesis of the fatty acid portion of the fat molecule is not yet clear. It is generally agreed that the first step in the synthesis involves the formation of reactive 2-carbon fragments, which are then condensed with each other. On the basis of available evidence, the reactive
2-carbon fragments arise by the breakdown of carbobydrate to the stage
of pyravic acid, as described above, followed by oxidative decarboxylatiou of the pyruvic acid. There is a strong possibility that acetyl CoA is
the reactive intermediate in the condensation, although the nature of this
condensation has not as yet been precisely defined. Much of the work on
fatty acid synthesis from earbohydrate has been done with bacterial
systems, and it is not certain to what extent the conclusions reached
are applicable to animal tissues. Although fatty acid synthesis has
been demonstrated in animal tissue preparations, these preparations
are usually too complex to permit a precise explanation of the reaction
mechanism.

The formation of neutral fat from carbohydrate by way of pyruvic acid would involve the production of reactive 2-carbon fragments, condensation, reduction of the \(\theta\)-keto compounds formed to saturated acids, and lastly esterification of the saturated acids. Acetoacetic acid itself is known not to be an intermediate in this synthesis, although the possibility remains that an active form of acetoacetic acid, e.g. acetoacety CoA, may be concerned. Acetic acid has been shown to be a precursor of the higher fatty acids, presumably after conversion to either acetyl-phosphate or acetyl CoA; fatty acid synthesis from labeled acetate bas

Masoro, Charloff, and Dauben: J. Bud. Chem., 179, 1117 (1949).

been demonstrated in liver homogenates * Moreover, the mammary gland has heen shown to be especially active in lipogenesis. Large amounts of acctate arise in ruminants, owing to the fermentation of carbohydrate by microorganisms in the rumen. This acctate is absorbed from the rumen by the blood and carried to the mammary gland, where it is used not only as a source of energy but also for fat synthesis, in fact, Popiák' reported that 50 per cent of the retained dose of acctate injected into a goat could he accounted for in the milk fat. The relationship of acctic acid to carbohydrate and fat metabolism is further discussed on p. 1009

The Blood Sugar. The sugar of the blood is glucose, present normally at a concentration of ahout 80 mg per 100 ml of blood, or 0 08 per cent This level is maintained constant within relatively narrow limits under all ordinary circumstances, significant variations from the normal range usually indicate some aberration of carhohydrate metaholism The con stancy of the hlood sugar level apparently is duo to a halance hetween the rate at which glucoso enters the blood and the rate at which it leaves the blood Blood glucose appears to originate chiefly in the liver, the rapid fall in blood glucose level after hepatectomy indicates that other tissues cannot contribute materially to the blood sugar content, although there is some evidence that the kidneys may play a minor role in this respect The glycogen of the liver usually is considered to ho the major source of the blood glucose, but the maintenance of a normal blood sugar lovel in the fasting animal long after liver glycogen stores are exhausted indicates that carhohydrate synthesis in the liver may also play an impor tant part in this connection

Factors which lead to a removal of glucose from the hlood include oxidation hy the tissues, conversion to glycogen in liver and muscle, and to a minor extent in other tissues storage as fat, and under certain conditions exerction in the unior. Increased oxidation in the tissues and increased conversion to liver and muscle glycogen each tend to lower the hlood sugar level when these two factors operate together, as after the administration of insulin, the resulting hypoglycema may be profound in the absence of insulin, as in diahetes mellitus the inability to store glycogen in the liver coupled with decreased oxidative utilization in the peripheral tissues generally is believed to be responsible for the elevated hlood sugar values seen in this condition. As alternate explanation for the hyperglycemia of diabetes mellitus is accelerated gluconeogeness ('overproduction') rather than impaired oxidative utilization ('underconsumption'') Evidence in favor of the overproduction theory is limited.

Increased Liveogenolysis in liver (but not in muscle), as after the administration of adrenaline or during ether anesthesia likewise causes an clevation of the blood sugar. The glycogen of muscle is not a direct factor in the maintenance of the blood sugar level, but operates only indirectly insofar as the muscles remove glucose for glycogen formation or contribute lactic acid to the blood which is theo carried to the liver,

Brady and Gurin Arch Biochem. Biophys 34, 221 (19o1)
Pop &k French and Folley Biochem. J 48 411 (19o1)

where blood lactate may be a significant source of liver glycogen and hence of blood glucose.

Storage of carhohydrate as fat (discussed on p. 996) is apparently a property of most of the tissues of the body. Loss of glucose from the blood because of excretion in the urine will occur whenever the rate of reahsorption of glucose hy the renal tuhular cells falls helind the rate of entrance of glucose into the glomerular ultrafiltrate-i.e., when the level of plasma glucose exceeds the renal threshold concentration. Ordinarily, the renal threshold concentration corresponds to a plasma glucose level of 150 mg. per cent or so, and this is only transitorily exceeded after ingestion of a large amount of carhohydrate, during emotional stress, etc. In the rare condition known as renal diahetes the renal reahsorption of glucose is so diminished that significant amounts escape into the urinc. and after poisoning with the drug phlorizin the renal reabsorption of glucose is almost completely aholished (phlorizin diabetes, see p. 1022). The exerction of blood glueose in the urine in diahetes mellitus is due more to the elevated concentration in the blood than to any significant alteration in the renal threshold. In none of the various conditions described which lead to excretiou of glucose in the urine is the loss from the blood ordinarily sufficient to affect appreciably the level of the blood sugar concentration.

Hormonal Control of Carbohydrate Metabolism. The metaholism of earhohydrate in the animal body is under the control of hormones from several of the endocrine glands, of which the pancreas, the anterior pituitary gland, and the adrenal cortex appear to be of major importance. Some aspects of this hormonal control have been indicated in the preceding pages; the endocrine glands themselves are discussed in Chapter 26.

Knowledge concerning the relation between the panereas and carbobydrate metaholism hegan with the classical demonstration by Von Moring and Minkowski in 1889 that removal of the pancreas in the dog was followed by development of the symptoms of diabetes mellitus. In 1921 Banting, Best, and Maeleod obtained the active principle of the pancreas in this connection, the hormone insulin. Administration of insulin to a normal animal leads to a profound hypoglycemia which may result in convulsions or unconsciousness (insulin shock). In the diabetic, a maintained administration of insulin will completely alleviate the condition. Opinions differ as to the mechanism of insulin action; some feel that insulin accelerates carbohydrate oxidation, while others regard its function as primarily concerned with the synthesis of carbohydrate from nonearbohydrate precursors (gluconeogenesis) or possibly with the conversion of carbohydrate to fat. The significance of insulin in the hevokinase reaction is discussed on the following page.

A further notable advance in knowledge concerning the hormonal control of carbohydrate metabolism was the demonstration by Houssay that removal of the pituitary gland ameliorated the symptoms of diabetes in the pancreatectomized dog. Removal of the adrenal cortex is likewise effective in ameliorating the symptoms of pancreatic diabetes (Long and Lukens). These and other results suggest that control of carbohydrate

metabolism by the hormones of the pancreas, the antenor pituitary gland, and the adrenal cortex is essentially a balance between opposing forces, with the effect of insulin opposed by either or both the pituitary and adrenal-cortex principles. Striking confirmation of this concept is afforded by the work of the Coris and their associates, who have found that the ability of tissue extracts to phosphorylate glucose by the hexokinase (glucokinase) reaction (see p. 990) is inhibited by a fraction of anterior-pituitary extract, and also by adrenal-cortical extract, this in hibition is overcome by insulin. This we extracts from diahetic animals showed an inhibited hexokinase activity which was overcome in ritro or in rivo by insulin. This was the first demonstration of the hormonal control of a specific enzymatic reaction of earbohydrate metabolism, these experiments have been confirmed by others.

In another series of experiments, Sutherland and Con¹º present evidence which indicates that the phosphorylase enzyme of the liver is under the control of epinephino and the hyperglycemic factor (HGF, see p 769) of the panereas, small amounts of epinephinie or of HGF markedly increased phosphorylase activity in liver slices where activity was low, and preserved activity in liver homogenates, similar effects were observed in nio It is suggested that phosphorylase activity in liver represents a balance between inactivation and resynthesis of active enzyme, and that thus is in some as yet unknown way under hormonal control. It will be recalled that hoth epincphinie and HGF promote the breakdown of liver

glyeogen in the animal hody

Knowledge concerning relationships between the various other hormones of the body and earbohydrate metaholism is relatively meager and difficult to evaluate The thyroid bormone influences the rate of metabolism in the tissues, but no specific relation to carbohydrate metabolism has been demonstrated, although a role in gluconcogenesis has been postulated A role in gluconcogenesis has likewise been delegated to certain of the steroid hormones of the adrenal cortex (see Chapter 26). The subject is rendered difficult by the lack of purity of the bormone preparations in some mistances and by the possibility that 4 postulated action or lack of action may in reality involve the mediation of some other gland or hormone in addition to the one thought to be concribed.

FAT METABOLISM

The fats of the diet and of the animal hody represent largely a concentrated form of energy, for metabolic or storage purposes, but there is adequate evidence that fats may subserve important nonealoric metabolic functions as well. In this connection, Burr and Barnes conclude that "there are ample reasons for recommending that the fat intake be not reduced much below the normal established by habit." Thus the fat contint of the diet may influence such diverse processes as the digestibility and absorbability of other foodstuffs in the gastrointestinal tract, and the rate of caleffication of the bones fatty acids represent important constituents of the lipids structural components of tissues, the fats of the

^{*} Sutl erland and Cert J Biol Chem 188 531 (1951)

diet serve as a vehicle for the fat-soluble vitamins, and the presence in the diet of certain highly unsaturated futty acids which cannot he svnthesized by the animal hody bas been shown to be necessary for normal growth and tissue metaholism Moreover, the presence of fat in proteinfree diets exerts a favorable effect on nitrogen retention, as well as on the course of pregnancy and lactation The physical capacity of rats has been shown to be greater on high fat dicts than on fat-free regimens Dietary fat has likewise heen shown to exert beneficial effects on rats subjected to such stress factors as thyrotoxicosis 11 x radiation, 16 and cold 13

The proportion of total calories furnished by dietary fat may vary widely from one group of individuals to another, and depends chiefly on such factors as availability, cost and established nutritional customs The recommendation of the National Research Council is that fat he present in the diet to the extent of furnishing 20 to 25 per cent of the total calories (30 to 35 per cent at higher levels of calorie expenditure) and that at least 1 per cent of the total calories he in the form of the essential unsaturated fatty acids (see p 1004)

Comparative Nutritive Value of Fats. The variety of fats found in nature has stimulated investigation into their relative nutritional merits for economic as well as for nutritive reasons These studies have failed to reveal any significant nutritional differences between the common animal and vegetable food fats and oils other than those attributable to slight variation in digestibility or in content of fat-soluble vitamins Deuel, et al 14 have furnished experimental evidence to refute the concept that hutter fat possesses certain saturated fatty acids not present in other fats, which are essential for growth. While rats prefer a diet containing hutter to one in which the fat is corn, cottonseed olive, peanut or soyhean oil or margarine this preference is apparently due to flavor alone mar garine fat and hutter fat promote similar growth responses when fed to weaning rats under controlled conditions 15 Deuel has reported that adequate growth and reproduction bave been maintained in rats through 36 generations on a diet in which skimmed milk powder and margarine fat replaced the whole milk powder of Sherman's well known whole milk powder-ground whole wheat hasal diet. The equivalence of margarine and butter diets in promoting the growth of children has been demonstrated in tests carried out over a two-year period under well controlled conditions 16 These and other results lead to the conclusion that butter fat possesses no special nutritive powers as compared to vegetable fats. and that properly fortified margarine is an adequate substitute for butter and has substantially equivalent nutritional value 17 Fortification of the first American margarine was carried out in the laboratory of the senior

Greenberg and Deuel J Vuirtion 42, 279 (19-0)
 Cleng hryder Berguust and Deuel J Vuirtio 48 161 (195)
 Mitceld Glackman La nbort Keeton and Fahmestock 1m J Physiol 146 S4

Deuel Movitt Hallman and Mattson J Nutrition 27 107 (1944)

Deuel Movitt and Hallman Serie ce 98 139 (1943)

Deuel Movitt and Hallman Serie ce 98 139 (1943)

Deuel Movitt Benberg, and Carlson J Am. Med. 1ssoc 136 388 (1948)

Deuel Science 133, 183 (1946) J Nutrition 32, 69 (1946) also Ld torul J Am. Wed 4ssoc 128 SS1 (1945)

author, who also demonstrated in rats that margarine so fortified was the nutritive equivalent of butter 15

Studies on the digestibility of natural fats and oils have shown that the common cooking oils such as cottonseed, corn, peanut, and obve oils, as well as butter, lard, and various margannes, are equally (95 per cent) digestible Beef and mutton tallow are somewhat less digestible Rapested oil, which contains a large proportion of crucic acid, has a digestibility coefficient of oily 82 in the rat, is although it is almost completely digested in man 'o Moreover, castor oil does not produce the catharism in guinea pigs, rabints, rats, or sheep that it does in human beings Under such conditions, it is utdized to the extent of 92 to 99 per cent il Among the fatty acids themselves, myristic and lauric acids are practically completely digestible by the rat when fed in olive oil, stearic acid has a low digestibility, and palmitic acid occupies an intermediate position il The triglycendes of these acids behave in the same manner

Relation Between Diet Fat and Body Fat. Prior to the work of Schoenhamer and Rittenberg on fat metabolism as studied with the aid of fats labeled with isotopic hydrogen (deuterium), it had been believed generally that the fats of the diet were directly metaholized and that only the excess above energy requirements was stored in the fat depots of the hody These authors" showed that even when fat is fed at a low level, some is first deposited in the tissues instead of being subjected to direct combustion The experimental details are of interest Normal mice were kept on a diet low in fat, and containing I per cent of linseed oil which had heen partially hydrogenated with deuterium Later post mortem examination of the body fat for the presence of the deuterio fat indicated that, even though the total amount of depot fat remained constant, 47 per cent of the dictary fat had been incorporated in the depot fat. Thus the animals were burning an approximately equal mixture of depot fat and food fat, in spite of the fact that the fat content of the diet was below that needed for energy purposes if completely hurned

Depot fat is therefore not mert storage maternal hut is constantly involved in metaholic processes. The admixture of dietary and tissue fat shown by such experiments as that just desembed is, bowever, subject to some modifying control, since the animal tends to produce a body fat of fairly uniform composition and succeeds moderately well as long as the fat of the diet is not altered to too great an extent If, however, large amounts of fats containing foreign types of fatty acids are taken in, some of these may be deposited in the tissues unchanged and may modify the character of the body fat Thus the melting point of the hody fat of the dog has been raised from the usual 20° C to 40° C by feeding mutton

tallow and has been decreased to 0° C by feeding linseed oil Anderson and Mendel fed rats diets of skimmed-milk powder and added fat, and found that the jodine number of the body fat could be varied from 122 to 35 by feeding soybean and eccount oils with iodine numbers of 132 and 77, respectively This matter is of commercial importance since feeding hogs rations too high in liquid fats gives rise to a soft body fat yielding a soft lard In such cases, feeding for some time of a diet high in carbohydrate has a hardening effect, since the fat formed from carbohydrate has a higher melting point and tends gradually to replace a portion of the lower-melting-point fats

However, animals vary considerably with regard to the susceptibility of tissue fats to alteration in composition due to the fats present in the dict Shorland24 designates the hody fats from species such as cows, sheep, and other ruminants which resist change due to diet as homolipides, while the fats from species which respond more or less readily to food fats of unusual composition by a change in body fat composition are called heterolimides In considering this problem from an evolutionary viewpoint, it has been suggested that the storage depot fats of the lower marine forms, which have the most complex fatty acid make-up of all species. merely mirror the composition of the food fats 25 This is because such ammals are unable to dilute the food fat with synthetic fats, since they do not possess the ability to change carbohydrate and protein into fat As the capacity for transforming these nonlinide components into fatty acids develops in the forms ranking higher in the evolutionary nattern. the modification of the storage fat by diet becomes less pronounced. In the ruminants, most of the storage fat is newly synthesized from protein and carbohydrate, dietary fat is largely modified in the complicated gastromtestmal tract, and it exerts no appreciable effect upon the composition of the fat in the storage depots According to Wertheimer and Shapiro, "6 synthesis of fat from carbohydrate actually takes place in the admose tissue

Some modification of ingested fat may occur during absorption and resynthesis in the intestinal wall. That the various saturated fatty acids found in the mixed triglycerides of animal body fats are in synthetic combrium is shown by the fact that, after the feeding of a particular fatty acid laheled with deuterium ie, deuterio-palmitic acid, a siginficant amount of the deuterium is subsequently found incorporated in the other saturated fatty acids of the body fats Deuterium is also found in the olcic and palmitoleic acid fractions, thus proving that these mono-unsaturated acids may be synthesized from dietary components "7 This synthesis may be by direct dehydrogenation, but it appears more probable that the fed fatty acid enters into degradation reactions to produce smaller fragments from which the oleic acid is then synthesized No deuterium is found in the linoleic reid fraction, thus confirming the original observations of Burr and Burr (see p 1004)

⁴ Shorlar 1 Nat re 165, 766 (1950)
25 Shorland Nature 1 0 924 (1352)
26 Wertleimer at 1 Stay tro Physiol Revs 28, 451 (1948) " Stetten an I Sel oenl eimer J Biol Chem 133 323 (1310)

that the animal is unable to synthesize these highly unsaturated fatty

Essential Fatty Acids The mahility of the animal hody to synthesize certain highly unsaturated fatty acids essential for its normal nutrition was first demonstrated by Burr and Burr,25 in 1929 These investigators found that rats placed on a diet devoid of fat hut otherwise apparently complete failed to grow and developed characteristic lesions of the skin and tail (see p 1073 and Fig 268 for complete description of this condition) Addition to the diet of the normal saturated fatty acids or of oleic acid did not render the diet complete, in fact, the administration of hydrogenated eccount oil has been shown to shorten the depletion period of rats on a dict free from essential fatty acids, as well as to intensify the accompanying symptoms 23 When triolein was the fat employed, this effect was not noted 30 When small amounts of highly unsaturated fatty acids such as linoleie or linoleine acid were added to the diet the deficiency did not develop Further study has confirmed these findings, the deficiency is apparently not concerned with the formation of fat from carbohydrate, or fat storage, arachidome acid is more effective than the other unsaturated fatty acids in promoting growth, and in the deficient annual the arachidome acid content of the liver is maintained at the expense of continuous depletion of the other tissues 11 This would imply that the presence of highly unsaturated fatty acids in liver, at one time thought to he a characteristic of fat metaholism in this organ, is in reality due to selection and retention of these compounds from the blood Arachidonic acid appears to be the essential fatty acid which is the physiological member of the group Using spectrophotometric procedures for the determination of the number of double honds in the fatty acid fraction of blood and tissue fats, Reiser12 and Witten and Holman 13 have demonstrated the synthesis of tetraenoie acid (arachidonic) from linoleic acid

Oxidation of Fats. The glycerol portion of the fat molecule is undonbtedly oxidized in the body by pathways of carbohydrate metabolism,
probably after preliminary pho-phorylation to form phosphoglycerol,
which then by oxidation could give rise to cither pho-phoglycerol addity de
or phosphoglycene acid, both of which are recognized intermediates in
carbohydrate breakdown (see p 989) The fatty acids are oxidized in
quite a different way

 β -extraction It is generally agreed that the major method whereby fatty acids are oxidized in the animal body is by the process known as β -excitation vecording to this concept, the fatty acid chain is oxidized at the carbon atom β to the carboxyl group, with the splitting off of a

2-carbon fragment and the production of a fatty acid baving two less carbon atoms than the original. This process continues along the chain until the entire fatty acid molecule has been hroken down to 2-carhon fragments by the removal of two carbon atoms at a time.

The first biological evidence for \$\beta\$-oxidation was afforded by the experiments of Knoop This investigator found that on feeding animals fatty acids of varying chain length hut in each case with the a-carhon (the one farthest removed from the carboxyl carbon) attached to a phenyl group, there could he isolated from the urine either henzoic acid or phenylacetic acid, in the form of the conjugated derivatives hippuric acid and phenaceturic acid, respectively. Benzoic acid was obtained when the fatty acid chain contained three or five carbon atoms; phenylacetic acid when it contained two (i.e. phenylacetic acid itself) or four carbon atoms. These results led Knoop to conclude that the carbon atoms could not be removed one by oue during oxidative breakdown of the fatty acid, hut must come off in pairs, i.e., by oxidative removal at the \$-carbon.

The mechanism of β-oxidation, and the nature aud fate of the 2-carbon fragment thereby produced, has been the subject of prolonged and intensive study. A major difficulty in the past, since overcome, has been the inability to obtain cell-free preparations from animal tissues which were capable of oxidizing fatty acids and were susceptible to experimental study. Muuoz and Leloir24 and Lehninger25 were the first to describe such preparations from liver tissue. Lehninger's preparations under the proper conditions readily brought about the oxidation of all of the normal saturated fatty acids containing from 4 to 16 carbon atoms, with acetoacetic acid as the main product. In the exidation of octanoic acid, for example, two moles of acetoacetic acid could be obtained for each mole of octanoic acid oxidized:

CH₃(CH₂)₄COOH + 2O₂ → 2CH₃·CO CH₂·COOH Octanoic acid Acetoacetic acid

The interesting observation was made that if oxalacetate was added to the system, the yield of acetoacetate decreased and some citrate, α-ketoglutarate, and succinate were formed, suggesting that the citric acid cycle of Krebs (see p. 990) was entering into the picture. Since added acetoacetate was not oxidized, even in the presence of oxalacetate, and pyruvate was found to be oxidized in a manner similar to the oxidation of fatty acids, it was postulated that both fatty acid oxidation and pyruvate oxidation produced a reactive 2-carbon fragment as a common intermediate. This 2-carbon compound could then condense either with itself to form acetoacetate as end product, or with oxalacetate to form citrate. the eitrate then entering the oxidative metabolism of the Krebs cycle. With the identification of the reactive 2-carbon fragment as the acetyl group of acetyl-coenzyme A (p. 1010), the process is now pictured by Lebninger as follows:

³⁴ Munoz and Leloir: J. Biol. Chem., 147, 355 (1943).

³⁵ Lehninger. J. Biol. Chem., 161, 437 (1945).

$$\begin{array}{c} \text{CH}_{2}(\text{CH}_{2})_{*}\text{COOH} \xrightarrow{\beta\text{-condat on}} \text{CH}_{2}\text{CO} \xrightarrow{S} \text{-CoA} \\ \text{Fatty acid} & \text{Acetyl CoA} \\ & \text{Acetyl CoA} \\ & \text{Acetoacetate} & \text{Citrate} \\ & \text{Krbs} \\ & \text{CO}_{2} + \text{H}_{2}\text{O} \end{array}$$

Further aspects of the metaholie significance of acetoacetic acid and acetyl groups are considered on pp 1007 and 1010 In connection with the breakdown of long chain fatty acids to the stage of acetyl groups it has been suggested that the first step in the reaction is the formation of a fatty acid-coenzyme \ combination, through the mediation of adenosinetripho phate (VTP) In the case of heart-muscle preparations which oxidize fatty acids, Vlahler" has formulated this reaction (for butyrate) as follows

where AMP is adenosine monophosphate, or adenytic acid. Presumably a similar reaction occurs with other fatty acids as well Evidence to this effect has been obtained by the use of hydroxylamine to trap" fatty acid esters of coenzyme 1 through the formation and identification of the corresponding hydroxamic acids

Using this procedure caprylyl hydroxamic acid has been isolated from liver extracts actively oxidizing caprylic acid the oxidation presumably going through the intermediate formation of a caprylyl-coenzyme A ester

A second method for the formation of fatty acid-coenzyme A com binations which thus far appears limited in its applicability to butyric and acctoacetic acids is through exchange with succinyl CoA According to Mahler 26 oxidation of α ketoglutaric acid to succinic acid in the presence of coenzyme 1 gives rise to the intermediate formation of succinyl Co \, which can then convert buty rie acid into butyryl Co \

It is to be noted that this reaction involves DPN (in the ketoglutaratesuccinate stage) but not ATP and it may account for the differences which have been noted occasionally between the oxidative metabolism of butyrie acid and that of other fatty acids.

It appears likely that the fatty acid-coenzyme 1 combination once formed is a substrate for a dehydrogenase action involving DPN and resulting in the formation of a 3-keto group in the fatty acid chain

^{**} Mahler I hosphorus Metabolum Vol II Baltim re J has Hopkins University Press.

¹²⁰

$$\begin{array}{l} \text{RCH-CH}_2\text{CO} \longrightarrow \text{S-CoA} + 2\text{DPV} \\ \rightarrow \text{RCOCH}_2\text{CO} \longrightarrow \text{S-CoA} + 2\text{DPNH} + 2\text{H}^+ \end{array}$$

According to current views, the β keto aeyl ester thus formed can react with another molecule of ecenzyme A at the keto group, to split off acetyl CoA and leave an acyl ecenzyme A combination containing two less carbon atoms

$$\begin{array}{c} \mathrm{RCOCH_2CO} - \mathrm{S-CoA} + \mathrm{HS-CoA} \\ \rightarrow \mathrm{RCO-S-CoA} + \mathrm{CH_3CO-S-CoA} \\ \end{array}$$

Thus by this thioclastic reaction the acetoacetyl Co \(\) formed in buty rate oxidation will give use to acetyl Co\(\), whereas for the higher fatty acids the combination of dehydrogenase activity and reaction with coenzyme \(A \) results in the shortening of the chain by \(2 \) carbon atoms at a time, with the concomitant release of acetyl groups in the form of acetyl Co\(\), until the entire molecule is broken down

As stated above, the acetyl groups which are the intermediate products of fatty acid oxidation give rise to acctoacetic acid or are further oxidized by the metabolic processes of the Lirebs cycle There is some evidence that acetyl groups derived from the carboxyl end of a fatty acid chain may bave a somewhat different metabolic significance from those derived from the metbyl end of the chain, further work will be needed to clarify this point There is much in the picture of fatty acid oxidation presented here that is obscure and that will presumably be subject to modification as knowledge increases in this field. At the same time it is believed that a reasonably satisfactory explanation is now available for most of the known facts regarding fatty acid ovidation in animal tissues and that this explanation represents a significant advance in understanding over previous concepts. It is likely that other pathways of fat oxidation exist. For example, it is known that fatty acids may undergo ω-oxidation—i e . oxidation beginning at the carbou atom fartbest removed from the carboxyl group-since certain long-chain dicarboxylic acids bave been isolated from the urine of animals after the feeding of fatty acids The significance, if any, of this type of oxidation in the normal metabolism of fatty acids is obscure

Metabolism of Acetoacetic Acid Acetoacetic acid, its equilibrium reduction product β-bydroxybutyne acid, CH₂ CHOH CH₂ COOH, and its decarboxylation product acetone, CH₂ CO CH₃ are the ketone bodies ('acetone bodies') which normally are found only in very small amounts in the blood and urine. Under certain conditions, as in diabetes mellitus, during starvation, or prolonged subsistence on a low carbohydrate duct, the amount of these ketone bodies in the blood rises and considerable quantities may be exerted in the urine (see experiment on p. 1073). Such a condition is known as ketosis. It was beheved for some time that acetoacutic acid and its associated ketone bodies were abnormal metabolic end products and in particular that acutoriche and represented the inability of the tissues to carry fatty acid oxidation by the β-oxidation process by youd the stage of the 1-carbon compound

It is now known that acetoreetic read is a normal and product of fatty and oxidation in liver. The acetoreetic acid formed in liver is not further

utilized by this organ, except possibly to a slight extent in fasting. Other tissues, however, readily metabolize acctoacetic acid to CO2 and II2O, and there appears to be no impairment in this respect in diabetes (Soskin) The intensity of ketone-body production (ketogenesis) by liver appears to be largely a question of substrate availability. If adequate earbohydrate is available, the hver apparently profers carbohydrate oxidation as a source of energy, and ketone-body production is small Carbohydrate is therefore an "antiketogenic" substance. In the absence of carbohydrate oxida tion, as in diabetes or when glycogen stores are exhausted, oxidative energy is derived almost entirely from fatty acid breakdown and ketone bodies result. They may be produced by the liver in such quantities that the peripheral tissues are unable to ovidize them as fast as they are formed, in which case they will accumulate in the blood (ketonemia) and he excreted in the urine (ketonuria)

It was felt at one time that ketogenesis was harmful and that it could be controlled by the proper ratio in the diet of ketogenic material (fats, and the ketogeme portion of proteins) to antiketogenic material (carboh) drate and the glucogenic portion of protein) These views are no longer held The major effect of ketosis on the animal body appears to be in rela tion to acid base balance, exerction of large amounts of acctoacetic acid and β-bydroxybutyric acid in the urine as their alkali salts depletes the body of available base and may lead to the development of an acidosis.

That ketone bodies may arise from sources other than fatty acid oxida tion is well recognized. Their origin from pyruvate under certain circumstances has been shown, and it is well established that certain amino acids such as leucine and phenylalanine are metabolized via the intermediate formation of acctoacctic acid Presumably any metahohe source of acetic acid could also serve as a source of acctoacetic acid (see p 1009) Thus ketone body production is not a characteristic of fatty acid oxidation alone, but is rather to be considered only one of the various metabolic processes which yield energy to the organism

Ketonuria during fasting occurs spontaneously to an appreciable extent only in man and the higher apes However when the salts of ketogenic acids are administered to fasting rats a marked ketonuria obtains, which is physiologic This is defined as an exogenous ketonuria Likewise, after the development of fatty livers in rats a spontaneous ketonuria of con siderable magnitude results on fasting this type of ketonuria is known as endogenous ketonuma Women exhibit a much higher degree of fasting ketonuria than do men " the same sex difference has been shown to occur in both the exogenous and the endogenous types of ketonuria in the rat

Further stages in the hreakdown of acetoacetic acid to CO2 and H2O occur largely in tissues other than the liver and are considered to involve the participation of coenzyme A and the reaction mechanism of the Krebs tricarboxylic acid cycle The role of the Krebs cycle in acetoacetate metabolism was first postulated by Breusch and by Wieland and has

[&]quot; Deuel and Gul ck J Biol Chem % 2. (1932)

"Butts and Deuel J Biol Chem 109 41. (1933)

" Deuel Hallman and Murray J Biol Chem 119 2.7 (1937),

" Breusel Science 97 4.00 (1943)

⁴¹ Wieland and Rosenthal Ann 554, 241 (1943)

been confirmed by other investigators. For example, Buebanan et al. 42 found that, after incubating kidney tissue with carbon-labeled acctoacetate in the presence of an excess of nonisotopic ketoglutarate, succurate, and fumarate, on subsequent isolation these three intermediates in the Krebs cycle contained isotopic carbon derived from the acctoacetate. These results, and those of many other investigators, leave no doubt concerning the participation of the Krebs cycle in acctoacetate mutabolism.

In the discussion of the \$\beta\$ oxidation of fatty acids on p 1005, it was pointed out that the production of acetoacetate itself as an intermediate in the breakdown of fatty acids by extrahepatic tissue is considered to be unlikely, rather, there is formed the acetoacetyl ester of coenzyme A. acctoacetyl CoA This 8-keto acyl ester can undergo the thioclastic leaction with more cochayme A described on p 1007, to give use to acetyl CoA, which then follows well recognized metabolic pathways discussed below If this view is correct, then acctoacetic acid formed by the hver and brought by the blood to the extrahepatic tissues must be converted into acctoacetyl CoA before being further metabolized. One mechanism for the activation of acetoacetic acid is found in the reactions described by Mahler36 and discussed earlier in this chapter, according to this report acctoacetate is similar to butyrate in its ability to react with succinyl CoA to give (presumably) acetoacetyl CoA and succinate If this or an equivalent reaction is found to prevail generally in the extrahepatic tissues, the metabolism of acetoacetate by these tissues becomes readily understood

The possible relationship between fat and carbohydrate metabolism exemplified by the entrance of the reactions of the tricarbovylic acid cyclo into acotoacetate metabolism may afford some basis for the aphorism that "fats burn in the fire of the carbobydrates," which has fallen into some disrepute in recent years, possibly because it is clearly not applicable to fat metabolism in the liver. Certainly if it can be shown that carbohydrate metabolism facilitates acetoacetate combustion (which has not as yet been done but which appears hickly) the existence of a "ketolytic" as well as an antiketogenic action of carbobydrate must once aguin he considered, at least in reference to organs other than liver. For further discussions of ketone bodies, see Chapters 29 and 31.

Metabolism of Acetic Acid Acetic and is the simplest possible fatty and with an even number of carbon atoms. It is readily metabolizable when fed to an ammal, or when meubated with liver, ladiner, or heart tissue (but not brain or muscle). It has been somewhat difficult to evaluate the metabolic significance of acetic acid in the past, because of its metabolic lability and the lack of precise methods for its quantitative determination. By the use of isotope labeling, acetic acid has been implicated in the synthesis of a variety of substances such as acetoacetic acid, fatty acids, cholesterol, protoporphyrin, glycogen, and the dicarboxylic amino acids as well as in the acetylation of choline to form acctylcholine and of such substances as sulfamilamide and p aminobenzoic acid, which are excreted in part as acetyl derivatives in the innic On the basis of the "dilution" of isotopic dictary acetate by the nomisotopic acetate of the tissues,

⁴² Buchanan Sakanu Gurin and Wilson J Biol Chem 159 630 (1945)

Bloch and Rittenberg⁴² estimate that a 100-g rat produces about 1 g of acetic acid per day by all metabolic processes, of which both carbolis drate breakdown and the \$-oxidation of fatty acids are major components Since the animal normally exerctes little if any acetic acid in the urine, all of the acctic acid produced must be further metabolized

The mechanisms of acetate metabolism have been intensively studied in many laboratories. As is evident from the previous discussions in this chapter, the oxidative breakdown of both carbohydrate and fatty acids appeared to require the formation of an intermediate 2-carbon compound which was clearly not acetate itself but which was closely similar to acetate, and hence called by many ochre acetate Studies on the generation of acctyl groups by animal tissues, e.g., the formation of acetylsul familamide from sulfamilamide by liver tissue, likewise pointed to the existence of an active form of acetate A major contribution to this field was made by Lipmann in the discovery of coelizyme A (for acctate) as the counzyme required in the enzymatic transformations of acetate by bacterial and aiumal tissues. Many illustrations of the role of coencyme 4 in carbohydrate and fat oxidation have already been presented in this chapter, the structure and vitamin relationships of coenzyme A will be found in Chapter 35

The search for active acetate appears to have reached a successful conclusion in the discovery by Lynen 4 of the formation of the S-acety ester of coenzyme A as an intermediate in the metabolism of acetate by yeast Coenzyme A contains a mercaptoethanolamine residue in the molecule, the relation between the structures of coenzyme A and acetyl eoenzyme A is as follows

Acctyl coenzyme A fulfills all of the requirements for active acetate in the test systems thus far studied For example, while ordinary acetate requires the presence of ATP to form citrate in tissue extracts containing oxalacetate, acetyl CoA yields citrate without ATP being present Simi larly, the requirement for a source of energy in the formation of acetyl eholine and acetylsulfamlamide by tissue preparations disappears if acetyl CoA rather than acetate is used as a source of acetyl groups Other examples of the significance of acetyl CoA in reactions requiring the presence of active acetate have been given in the discussions of carbohydrate and fat oxidation

It is clear therefore that acetate as such is essentially mert metaboli cally until it is activated by conversion to acetyl CoA That such activa tion occurs in animal tissues is generally believed, the mechanism how ever is still not clear Lipmann has shown in both yeast and liver extracts that the formation of acctyl Co I from acctate in the presence of ATP may occur as follows

⁴ Bloch and Rittenberg J Biol Chem 159 45 (194.)

[&]quot;Lynen and Ruckert Angew Chem 63 4" (13-1)
"Lynen and Jones Black at I Flysis J im Chem Soc 74 2384 (1972)

The formation of acetyl derivatives of foreign aromatic amines (sul fanilamide, p-aminoberzote acid) and amino acids (phenylaminobutyric acid) has been studied with the aid of the various available isotopes, and by other means, both acetic acid and py rive acid have been shown to be precursors of acetyl groups in this connection, with possible qualitative and quantitative differences between them. The significance of the origin of acetyl amino acids from pyrivate is further discussed on p. 1018, in connection with amino acid metabolism.

Interconvertibility of Fat and Carbohydrate. That the fat of the hody may arise from earhohydrate has long been known (see p 996) The reverse process, the production of earbohy drate from fat and particularly from the fatty acid portion, has been the subject of considerable controversy Experimental methods based upon the production of extra glucose or gly cogen by which it is readily possible to demonstrate the glucogenic power of certain amino acids, for example, yield essentially negative results when fatty acids are studied For these and other reasons, many have maintained that fats do not give rise to earbohy drate in the animal body, although this conversion can be demonstrated in certain other organisms The opposing view, held by some, is that gluconcogenesis from fatty acids is readily possible and indeed is the major explanation for the hyperglycemia of diabetes mellitus. Much of the evidence cited in favor of this latter view is faulty. It has nevertheless been unequivocally demonstrated by the use of isotopes that the carbon of certain fatty acids can be incorporated into the gly cogen of the animal body, " thus proving that fatty acids and carbohy drate are reversibly related, as would indeed be expected from the equilibrium relations shown on p 1011 The amount of conversion found, however, is extremely small relative to that expected if the conversion of fatty acid to carbohydrate were of any value to the organism It has been suggested that such conversion must be uncconomi cal to the body in the light of current concepts of carbohy drate metabolism, since carbohy drate formation from fatty acids must inevitably be at the expense of the metabolic energy of carboby drate hreakdown Whether or not the altered metabobe state of the diahetic facilitates or even requires the conversion of fatty acid to carboby drate still remains to be unequivocally demonstrated.

Lipotropic Factors. On diets which are bigh in fats containing much saturated fatty and and which are low in protein or in choline, there is observed a large increase in the fat content of the line r. A high chole-terol diet likewise leads to the production of a fatty liver, the lipides in this case consisting of cholesterol as well as neutral fat. In either case, an increase in choline in the diet hrings about a reduction in liver fat. This is of interest in connection with the possible role of lecithin in fat metaholism, but the explanation for the so-called lipotropic action of choline is not yet established. Methionine likewise exerts a marked lipotropic action owing to its ability to promote the synthesis of choline by the transfer of methyl groups to suitable precursors (see p. 1024).

[&]quot;Buchanan Hastings and Nesbett J Biol Chem 150 413 (1943) Lorber Lifeon and Wood J Biol Chem. 161 411 (1945)

Other substances which evert a hpotropic action include lipocaic, a substance of as yet unknown nature claimed to he present in the panereas, mositol, a member of the vitamin B complex (Chapter 35) and a constituent of certain phospholipides (Chapter 11), tryptophan, and possibly glutamie acid Whether these various substances act in an independent fashion or through the mediation of choline action is not clear Tatty livers are also found under certain conditions in animals deficient in the essential fatty acids, in pantothenic acid or in rihoflavin, or in animals which have been provided with an excess of thiamine or biotin (Chapter 35) The relation between these various dietary constituents and the denosition of liver fat is not well defined, it is felt by some that they may represent nonspecific factors acting through some general change in the nutritional state

Parenteral Fat Administration. Although glucose and amino acids have been employed successfully in intravenous therapy for a number of years, practical procedures for the administration of fat are still in the process of development The two obstacles which prevented the earlier application of the procedure to man were the difficulty of ohtaining a preparation stable enough to resist sterilization and which would not cruse fat embolism. However, as early as 1935, in this country, Holt and associates.47 employing a butter emulsion stabilized with purified commer cial egg lecithin and containing 7 to 75 per cent of lipides, administered the preparation successfully to infants. The emulsion was prepared by homogenization at 4000 pounds pressure, followed by sterilization, most of the particles were less than 60 µ in diameter. Since this earlier work, a number of satisfactory fat emulsions have been prepared, using hutterfat, corn oil, ecconut oil, or other fats, hy employing soya phosphatide,45 or glycerol monostearate49 as a stabilizer, or by subjecting the fat to supersome radiations 50 Shafiroff et al 51 reported that, when fat was administered subcutaneously, a hetter utilization obtained when the spreading agent hyaluronidase was incorporated in the emulsion

Proof that emulsified fats are utilized following their intravenous administration has been obtained by the use of tagged compounds 52 For example, Lerner and eo-workers have found that, when a C14 labeled palmitie acid was injected intravenously into rats as a tripalmitin emulsion, as much as 59 per cent of the administered isotopic earbon was expired as CO2 within 24 hours, moreover, considerable amounts of the injected fatty acid were stored in the several adipose tissues. The bulk of the C14-labeled fatty acid stored in the liver and intestine was present in the phospholipide fraction Further confirmation that fat can be used when

⁴ Holt Tidwell and Scott J Pediat 6 151 (1935)

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"Upera and Blumberg Prec Sec Exp Biol Ved. 25 79 (1931)

"Shafiroff Baron Reelt and Vulholland Prec Sec Exp Biol Ved. 77 (508 (13-1))

"Shafiroff Baron Reelt and Vulholland Prec Sec Exp Biol Ved. 25 1110 (1949) Lerner Claikoff

Latenman and Dauber Prec Sec Exp Biol Ved. 25 (1949) 31 affiroff Vulhollar I and Baker hap Wed and Surg 9, 180 (19.1)

injected intravenously has been adduced by Mann et al, 52 who proved that it spares protein in growing pupples, and provides energy, as much as 30 per cuit of the total energy requirement could be used efficiently in the form of intravenous fat

Gly cerides of short-chain acids were reported to be toxic, while those of lauric, myristic, palmitic, and steame acids were found to be nontone when injected intravenously as emulsions in man Olcic and linoleic acid esters can be tolerated by human subjects, at least in small concentra tions 54 Geyer and associates55 reported a moderate increase in the total lipides in the spleen, lungs, and liver when stable emulsions of fats were administered by vein to rats Gorens et al - successfully administered as much as 3 g of fat daily per kilogram of body weight to adults and 6 g of fat per kilogram of body weight to a 7-week-old infant for periods up to 27 days Under these conditions weight loss was prevented and po time nitrogen and potassium halances were maintained Furthermore, fat injections produced no gross or microscopic changes in the tissues How ever, one should not overlook the report that intravenous fat produces a febrile response, which is termed thermogenic rather than pyrogenic, since it is not of hacterial origin 57 Lambert and associates 57 suggest that this effect may possibly result from an overburdening of the fat storage facilities due to the rapid influx of this foodstuff into the tissues Considerable work is in progress to find methods of preparation of fat emul sions which will not produce the febrile reaction

PROTEIN METABOLISM

The proteins of the diet are considered to be completely broken down to their constituent amino acids in the digestive tract by the action of the various proteolytic eozymes present and to be absorbed into the animal body in the form of these individual amino acids. The requirement for protem is therefore fundamentally (if perhaps not exclusively) a requirement for amino acids and it should be possible to express the protein requirement of an animal in terms of the amount and kind of amino acids rather than of protein itself. The ability of amino acids to replace protein in the animal diet was first demonstrated many years ago by the use of hydrolyzed protein supplemented with those amino acids known to be lost during the hydrolysis in place of protein itself to supply the nitrogen requirements of the animal This demonstration reaches its peak in the experimental procedure available today for both human and animal ex periments whereby mixtures of highly purified amino acids, in many cases synthetic products are used to replace protein completely in expen mental diets. The availability of this type of experimental procedure which was achieved only after the isolation and characterization of the amino acid threonine by McCov Mixer and Rose in 1935, has led to

major advances in the science of nutrition. It is also largely responsible for directing attention to the clinical practicality of supplying nitrogen requirements during disease and convalescence by either feeding concentrated intact proteins or intravenous administration of hydrolyzates and amino acid mixtures, an application of nutritional science which promises to be of outstanding medical value. It is an interesting commentary on scientific progress that approximately four decades elapsed between the demonstration that protein hydrolyzates were of nutritional value and the extensive application of this fact in medical practice.

Dispensable and Indispensable Amino Acids. The amino acids supplied by the protein of the diet are needed for the synthesis of new body proteins during growth and for the continuous regeneration of the tissue proteins of the adult, as well as for many special purposes such as the formation of hormones, enzymes, purines, hile salts, creatine, and many other similar compounds. It has long been recognized that certain of the amino acids needed by the animal body for these various purposes need not necessarily be present in the proteins of the diet, since they can be synthesized within the tissues from suitable precursors. On the other hand, there are certain amine acids required by the hody which cannot be synthesized under ordinary circumstances and which therefore must be present in the diet in adequate amount or nutritional failure, leading ultimately to death, will result.

Those amino acids which can be synthesized by the body from the other constituents of the ordinary diet at a rate adequate for nutritional demands are called dispensable or nonessential amino acids; those which cannot be so synthesized but must be present in the diet are called indispensable or essential amino acids. It must be clearly understood that these terms refer solely to the presence of these amino acids in the diet; as far as we know, all of the amino acid constituents of the protein molecule are essential in one way or another for the metabolic processes of the animal body. In fact, as du Vigueaud has pointed out, the so-called nonessential amino acids are more properly regarded as being so essential to the animal economy that it has been forced to retain the ability to synthesize them even at the expense of other constituents of the diet.

Knowledge concerning the dietary dispensability or indispensability of various amino acids has been obtained in the past largely by feeding experiments with young animals (e.g., rats) using a diet containing protein known to be low or lacking in certain specific amino acids, or containing a protein hydrolyzate from which certain amino acids (e.g., tyrosine, tryptophan) could be removed. Failure of the animal to grow on such a diet, followed by good growth when the diet was supplemented by the missing amino acids, afforded evidence as to the indispensability of the amino acids in question.

This procedure has certain obvious limitations and occasionally gave obscure results. A more adequate experimental basis is afforded by the use of mixtures of pure amino acids as the sole source of dietary nitrogen (Rose). To establish the nutritional significance of any one amino acid, it is only necessary to prepare the mixture without that particular amino acid and to use this deficient mixture as the source of nitrogen for the animal Nuch progress has been made in this field since this experimental technique became available

As a result of the extensive experiments of Osborne and Mendel, of Hopkins and of Rose, using the various procedures just described, the following tentative classification of the common amino acids with regard to their dictiry dispensability or indispensability for the young growing mainmal was suggested by Block

Indispensal le	ealle Partis Indispensable		Dispensable	
Histidine I j sine Tryptophan Pf en Jalanine Methic nine Threonine Leucine Isoleucine Valine	Group 1 Cystine Tyrosine	Group B Arginine Clycine	Clutamic seid Aspartic seid Alanine Serine Proline Hydroxy proline	

As one might expect, the division between the various groups of amino acids is not entirely a sharp one. Thus although methionine in sufficient amount will supply the needs of the animal for cystine, cystine is an important constituent of the detr in that it is able to spare that portion of methionine which otherwise would be converted into cystine. I similar situation holds true with respect to the conversion of phenylalanine into tyrosine.

The amino acids in Group B, arginine and glycine, apparently can be synthesized by young mammals but often at a rate insufficient to permit maximum growth. On the other hand hoth of these amino acids are necessary for normal growth of chicks This is an instance where an amino acid may be dispensable for one species of animal and indispensable for another

Knowledge concerning the amino acid requirements of man is somewhat limited. Of the amino acids listed above as indispensable for the young animal, histidine is apparently dispensable from the det of the adult human, at least as judged by the enterion of maintenance of introgen equilibrium (see p. 1020). Its quite probable that as more specific enterial than growth or nitrogen balance are used a somewhat different picture will be obtained than that described here. The conservative policy in human nutrition would indicate the presence in the diet of adequate amounts of all the amino acids listed in the first three columns above, even though experiments covering a short period tested.

Three suggestions concerning the recommended daily intake of indispensable amino acids for human nutrition have been made. One is calcu-

For review see Melmck J Am Dielet. Assoc 19 68a (1943)
 Rose Hames, and Warner J Biol Chem. 284 421 (1354)

lated from Rose's data on twice the minimum daily intake of each amino acid required to induce nitrogen equilibrium in young healthy adult men, another by Block and Bolling, calculated from the estimated per capita consumption of the principal protein foods in the United States in 1944, and the third from experiments on young soldiers on the low calorie (900 calones per day) army survival ration 60 These suggestions are summanzed below

RECOMMENDED DAILY CONSUMPTION OF INDISPENSABLE ANIXO ACIDS (AVERAGE FOR ENTIRE POPULATION)

Amino leid	From Rose ⁴¹ (g per day)	Block an l Bolling (g per day)	US Army Survival Ration (g per da j)
Arginine		4 8	3 1
Histidine	!	2 1	07
Lysine	16	4.7	3 3
Tryptophan	0.5	11	10
Phenylalanme	2 2	50	5 5*
Methionine	2 2	3 6t	3 St
Threonine	10	3 4	2 6
Leucino	2 2	83	4 2
Isoleucine	14	5 3	3 5
Valine	16	5 3	4 3

^{*} Includes tyrosine † Includes cystine

Origin of Amino Acids Although plants are able to synthesize all the known amino acids when supplied with a source of nitrogen, mor game as well as organic, the higher animals at least are able to synthesize only a little more than one-half of the common amino acids from the ordinary constituents of the diet or tissues. The limiting factor in the case of those amino acids which cannot ordinarily be synthesized appears in many instances to be the carbon chain or ring structure and not the nitrogen, since it is frequently found experimentally that the synthetic keto- or hydroxy-acid analog of the amino acid will take the place of the latter in the dict

R	R	R
CH ₂	ÇH₂	Ċн.
HCNH ₂	çо	нсон
COOH Amino acid	COOH Keto acid	COOH

⁶⁸ St ceter Vulnum Pers 10 289 (1959)
11 It is a t believed that the avalues which are only twice the minimum daily requirements of young lealth; adult makes at oil be used as He recorn ended amino act in takes of the entire \merican population

Sometimes but not always both keto and and hydroxy and are effective in this respect the hydroxy and presumably gives rise to the keto and by oxidation. Replacement of a dictary amino and by its corresponding keto and indicates that metabolism of the amino and proceeds reversibly through the keto and stage.

Synthesis of an amino acid from a keto acid usually is considered to involve a reversible equalit min with animonia this may proceed through the intermediate formation of the imino acid

$$R CH_2 CH(\ \ H_2) COOH \xrightarrow{+2H} R CH_2 C \ \ H COOH \xrightarrow{-H_2O} + H_3O$$

Amino acid

The biological formation of the amino acid from the keto acid may also in volve acetylation (du Vigneaud and Irish) *2

Another mechanism for the formation of amino acid from acto acid is by the transamination reactions described on p 1021 1t is not possible at the present time to evaluate the relative significance of these various processes in the synthesis of maino acids within tissues

The amino acids found in animal tissues are all of the \mathbf{L} configuration (see Chapter 4). In the case of certain amino acids it has been found that the unnatural ' or \mathbf{r} configuration is convertible into the natural form by the body. This presumably involves loss of the asymmetry around the α carbon by conversion of the amino acid to the keto acid followed by asymmetric synthesis of the amino acid in its natural configuration. Enzymes capable of converting \mathbf{r} amino acids to keto acids are found in many animal tissues. In supplying the amino acid needs of an animal of human with the synthetic \mathbf{r} amino acid however it is usually assumed that the \mathbf{r} component will be unavailable and twice as much of the \mathbf{r} mixture is supplied as compared to the requirement for the \mathbf{r} form for some amino acids the \mathbf{r} form is largely exercted in the urine unchanged but there are instain as (e.g. serie) where the \mathbf{r} amino acid has been shown to pr duce them aminostations

 $^{^{\}circ}$ lu V gr aud an I l isl J B of then 122 34) (1337) See also Bloch and Bo ek f Biol Chem 164 483 (1940

Daily Protein Requirement. The amount of protein required by the individual per day is a nutritional factor of obvious practical significance. A variety of conditions will clearly influence estimations of the protein requirement. Among these may be mentioned growth, pregnaucy, or other special demands of the iudividual; digestibility and absorbability and amino acid composition of the protein; possibly economic availability, individual idiosyncrasy, etc. Estimates of the protein requirement of mau usually have been based on studies of the nutritional status of groups of individuals on varying kinds and amounts of protein, and on the extension of the results of animal feeding experiments to man. The Food and Nutrition Board of the National Research Council has recommended that the protein intake of an adult be 1.0 g. per kg. per day, of good quality protein, for adequate nutrition. This corresponds to about 65 g. per day for an average adult human, and will result in a daily urinary nitrogen excretion of about 10 g. The recommended protein allowance for the female is increased during pregnancy and lactation; and for children it varies with age.63

The efficiency with which a given protein supplies the nitrogen requirements of an animal may be defined in terms of its so-called "biological value." Although the methods used in the past for the determination of biological value may be subject to revision in the light of changing concepts of protein metabolism (see p. 1048), the concept itself retains its usefulness. Broadly speaking, a protein of high biological value is one which bas a high digestibility and absorbability and supplies the organism with adequate amounts of those amino acids which it needs. The amino acids will include not only those which cannot be synthesized by the animal, but also sufficient of the dispensable amino acids to minimize the requirements for their synthesis. The presence of low or inadequate amounts of even one of the indispensable amino acids may be the limiting factor in the biological value of a protein. For example, if one such amino acid were present in a protein in such low amount that at ordinary levels of ingestion of the protein only half the animal's requirement for this amino acid were met, a untritional deficiency would result unless the protein intake were raised considerably above the usual level, thus leading to a lowered efficiency with respect to utilization of the other amino acids present. In general, animal proteins are found to have a higher hiological value than plant proteins hecause of the more satisfactory distribution in kind and amount of the various amino acids present. Endosperm proteins of the cereal grains (corn, wheat, rice) are lower in certain of the indispensable aming acids, particularly lysine, than are the commonly consumed proteins of animal origin. This has led to the erroneous generalization that all plant proteins are incomplete or poorly halanced with respect to their essential amino acid composition; as a matter of fact, many vegetable proteins, such as those of oats, heans, yeasts, wheat and corn embryos, leafy vegetables, and grasses, are almost as suitable sources of the amino acids as many of the more expensive animal products. It is sometimes recommended that at least half the protein of the diet be of animal origin. This

⁴² See table, p. 1108.

is not necessarily an adequate criterion per se, and several proteins of low biological value individually may so complement one another with regard to amino acid composition as to provide a mixture of high biological value

provided that they are ingested simultaneously

Since the absence of any one of the indispensable amino acids will result in incomplete retention (utilization) of the remainder, it has been proposed64 that the biological value of certain proteins may be related not only to their content of indispensable amino acids but to the relative rates of their release in and absorption from the intestinal tract For example, the biological value of soy protein is increased when the raw soybean meal is autoclaved, while that of easein is diminished by baking in neither case is any difference in indispensable amino acid content observed by analysis Nevertheless it has been demonstrated that the processing of soy protein increases the rates of release of methionine leucine, and lysine during enzymatic digestion but at relatively different rates The effect of differential rates of absorption in rito would be to furnish incomplete mixtures of absorbed amino acids at the early stage of absorption followed by the too late release and absorption of the supplementary amino acids needed for most efficient utilization 1t 15 well known that ingested amino acids do not accumulate in the blood stream Hence the proteins of highest biological value are those that not only contain the essential amino acids in adequate amounts but make them available for absorption at relative rates consistent with most efficient utilization for protein synthesis and retention

Nitrogen Balance and Nitrogen Equilibrium. The relation between the amount of nitrogen entering the hody from the diet in the form of ammo acids, and the amount of nitrogen excreted from the body in the form of metabolic end products (chiefly as urea but to some extent as unc acid, treatinine, etc) is known as the nitrogen balance. The nitrogen balance is positive if intake exceeds output negative if output exceeds in take, and if intake and output are essentially equal mitrogen equilibrium results The young growing animal must be in positive mtrogen balance since a certain portion of the ingested nitrogen is retained as newly formed body proteins and nonprotein nitrogenous compounds. During fasting or illness associated with wastage of tissues, a negative mirogen balance exists the normal healthy adult is ordinarily in a state of mirrogen equilibrium That is if the dietary intake of nitrogen is say, 15 g per day the total nitrogen excreted by all channels (urine feces skin) will be approximately 15 g. Of this excreted nitrogen, the urine ordinarily con tributes about 90 per cent

Sitrogen equilibrium may be established at almost any desired level of mtrogen intake from as low as 2 g per day to 25 or 30 g or even more. This apparently is because within these limits the intensity of mirrogen metabolism is determined by the rate of entrance of nitrogen into the body It was thought at one time that equality of nitrogen intake and ex erction represented essentially a disposal of dietary nitrogen in excess of the needs of the animal that is, a small portion of the entering mtrogen

⁴⁴ Vielnick Oser and Weiss Science 183 326 (1946)

was considered to be utilized by the hody for the replacement of utrogen loss due to wear and tear on the hody tissues, but the bulk of dietary uitrogen was essentially surplus material and was promptly utilized for energy-yielding purposes without contributing to the nitrogen metabolism of the tissues. This was the basis for the elassical distinction of Folin between endogenous and exogenous nitrogen metabolism; according to this concept, the bulk of excreted nitrogen was of exogenous origin and did not arise by the metabolism of tissue protein.

It is now known that this concept is not true, and that in general the nitrogenous constituents of the diet promptly enter into the varied mtrogen and protein metabolism of the hody tissues, becoming indistinguishable from similar substances already present, and the equality between nitrogen intake and output is due primarily to the fact that the rate at which a certain amino acid is metabolized is determined largely by the rate at which that amino acid hecomes available to the tissues. In other words, the entrance of a given amount of amino acid into the mctabolic processes of the tissue brings about the metabolism of an equal amount of the amino acid molecules which are already present. Thus the distinction hetween endogenous and evogenous hreakdown disappears, and must be replaced by the concept of a continuous and dynamic nitrogen metabolism the rate of which is determined, as are all chemical reactions, by the coucentration of reactants present at a particular time. The only biological exception to this thus far discovered is in connection with purine metabolism; apparently dietary purmes cannot penetrate the cell nucleus and thus enter into the endogenous metabolism of nuclear purines. 65

This overthrow of the classical distinction between endogenous and exogenous metaholism is due largely to the pioneer work of Schoenheimer and his associates, based upon the application of isotopes to biological problems. The details of some of this work arc instructive. Various amino acids (e g , leucine, glycine) were synthesized in the presence of isotopic nitrogeu (N15) so that the amino acid contained a significant amount of the isotope. These amino acids were then incorporated in small amount in the diet of rats. The urine was collected over a three-day period, after which the animals were sacrificed and the body nitrogen fractionated into protein and nonprotein portions. These, as well as the excreta, were then analyzed for the presence of the isotopic nitrogen. According to the classical concept of endogenous and exogenous metabolism, the urinary urea should have contained most of the dietary nitrogen and therefore most of the isotopic nitrogen should have appeared promptly in the urine. This did not happen. Less than one-half of the isotopic nitrogen of the glycine was excreted, and less than one-third of that of the leucine. The bulk of the unexcreted isotopic mtrogen was found in the tissue proteins. Later experiments have shown that the labeled amino acid is to a certain extent incorporated directly into the tissue proteins, and likewise contributes its nitrogen to various other amino acids of the body, since both the fed amino acid and other amino acids isolated from the tissue proteins were proved to contain the isotope.

[&]quot; Plentl and Schoenheimer: J. Biol Chem., 153, 203 (1944).

In general therefore one may state that the amino acids of the detenter rapidly into biological equilibrium with the amino acids of the body becoming incorporated into newly formed protein or entering into reactions which supply introgen for the synthesis of other amino acids or other introgenous constituents of the tissues. These reactions occur more or less rapidly among the various tissues and somewhat independently of the nutritional state of the animal, labeled amino acids are found incorporated into the animal body proteins both when there is an abundance of that particular amino acid in the diet, and when on a nitrogen free diet the tissue proteins are heing extensively broken down for energy purposes.

Conversion of Protein to Carbohydrate and Fat It has been well established that after metabolic removal of introgen, the carbon chain of certain of the anima located may be utilized by the animal for the formation of carbohydrate. The classical basis for demonstrating this formation entails the use of the drug phlorizm. If an animal is treated with phlorizm the renal threshold for glueose is lowered to such a degree that an immalistered glueose, or that formed within the animal body by metabolic processes, is excreted almost quantitatively in the urine ("phlorizm diabetes"). A fasting phlorizmized animal will continue to excrete glueose in the urine long after all carbohydrate stores have heen exhausted It is usually (but not invariably) found that such an animal excretes about 36 g of glueose for every gram of urinary nitrogen. The ratio of urinary glueose (dextrose) to urinary nitrogen is known as the D/\ ratio, which in this instance is 36.

A D/\ ratio of 3 6 usually is interpreted to mean that out of every 100 g of body protein metabolized by the fasting animal which would correspond to the exerction of 16 g of urnary nitrogen about 38 g of glucose are formed since 2\(^3\)\(_6 = 3\

If to a fasting phlorizinized animal either protein or certain amino acids are administred extra glucose is found in the urine along with extra inctabolic introgen. By quantitative measurement it is believed possible to evaluate the ability of either protein or amino acids to yield glucose in the aminal hody. Using this method, it has been found for example that various proteins yield from about 00 to as high as 80 per cent of their weight as glucose and that certain amino acids are glucose-formers while others are not. Those amino acids which have been shown to be glucogeneby this procedure include glycine alanine cystaine methionine nor fucione proline senne value arginine aspartic acid and glutame acid. Those amino acids which do not yield extra glucose in the phlorizing animal include isolaucine leucine lysine phenylalanine tyrosine and tryptophan.

The allihty of protein to yield glucose would thus appear to be related to the relative proportion of glucogenic amino acids in the molecule and wide variation among proteins may be expected. It is interesting to note that there is a rough parallelism between the glucogenic amino acids and

those capable of being synthesized within the animal body. In the light of current knowledge concerning intermediary carbohydrate and protein metabolism, it is not surprising that such amino acids as alanine, glutamic acid, and aspartic acid are glucogenie, since after metabolic removal of their introgen there remain the substances pyrivic acid, ketoglutaric acid, and oxilacetic acid respectively, and these latter compounds are recognized intermediates in carbohydrate hreakdown and synthesis.

Other methods of studying the conversion of amino acids to carbohydrate include the use of perfusion through isolated organs, the alibity of the compound to lead to increased liver glycogen content when administered to the fasting rat, and the ability of the amino acid to reduce an experimentally induced ketosis. Results by the various methods described are not always concordant for a particular amino acid. Tryptophan, for example, will reduce an experimental ketosis but will not lead to an increased liver glycogen content or an increased exertion of glucose in the phlorizinized dog, and other examples might be cited. It is possible that these various procedures measure metaholically independent functions rather than the same general property of carbohydrate formation, but more must be learned about the judividual reactions concerned before the subject will be better understood.

The couversion of amino acids to fats or fat metabolites is less well understood than that of carbohy drate formation. Certain of the amino acids (e.g., tyrosine, phenylalanine, leucine) appear to he oxidized directly by way of the formation of acetoacetic acid, and this may in turn give rise to the synthesis of fatty acids. Since proteins can be converted into glucose, and glucose into fat, some fat may arise from protein in this fashion. The significance of this in normal nutrition is uncertain.

Protein Storage. There is as yet no evidence for the existence within the animal hody of a storage form of protein analogous to the storage of carbolydrate as glycogeu, or of fat Yet there does appear to he a reserve protein supply which can he drawn upon to furnish the fundamental autrogeu requirements of the animal when the protein intake is madequate. This reserve protein appears to he drawn from the tissues themselves, such organs as the liver and kidney, and the blood plasma, appear to be capable of undergoing a considerable depletion of protein content to supply the needs of other parts of the body for nitrogen during fasting Other organs—e.g., hrain—are more resistant to protein depletion

Specific Dynamic Action of Protein. When protein is fed, more heat is produced in the hody than can be accounted for by the combustion of the protein ingested. One view is that this action is due to the stimulating action on the tissue cells of certain products of amino acid catabolism, perhaps hydrovy acids, leading to increased oxidation of carbohydrate by such cells. Glycine, alamine, and phenylalamine appear responsible for much of the specific dynamic action of proteins. Carbohydrates and fats also evert a specific dynamic action, but to a lesser extent than proteins and minio acids. For a further discussion of this subject, the reader is referred to Borsook's and kinss.

[&]quot; Borsook Biol Revs 11, 147 (1936)
" briss J Vutration 21, 257 (1941)

General Physiological Transformation of Amino Acids The amino acids in the body undergo many varied and complicated reactions. Some of these are being revealed in detail as a result of the extensive use of isotope techniques

TRANSIMINATION This refers to the transfer of amino groups from one compound to another. An amino and under the influence of a specific enzyme loses its amino group to a keto and and the original compound becomes a keto and

The keto acids found in these reversible reactions are concerned in oxidation reduction systems (carbobydrate metabolism) and thus may furnish amino acids or carbobydrate intermediates as required, 1e transamination is one link between protein and carbobydrate intermediary metabolism

Transmeritzation When methyl groups are linked to the nitrogen or sulfur of certain organic compounds they are capable of being shifted intact among these substances (du Vizneaud)

NH₂ Methlonine

Ethanolamine

These so-called labile methyl groups may then react with other labile methyl troup substances (guandoacetic acid, meeting acid, punnes etc.) to methylate these compounds

$$CH_i^- + \lambda H - CH_1COOH \rightarrow CH_1\lambda - CH_2COOH$$

$$C = \lambda H$$

$$\lambda H_1$$

$$Cuanidoacetic acid$$

$$Creating$$

⁵⁶ See p 1029

However, the methyl groups now become fixed and are no longer able to transmethylate. Further aspects of transamination and transmethylation are given on p. 1034 and p. 1029. Reactions thought to be concerned are given here:

METABOLISM OF INDIVIDUAL AMINO ACIDS

Glycine. This amino acid enters into a variety of metabolic functions. In addition to its presence in many of the body proteins (collagen, for example, is rich in glycine), it is concerned in the synthesis of such non-protein compounds as (1) the glycocholic acid of the liver, (2) the creatine of muscle and other tissues, (3) the glutathione of the cells, (4) the protoprophyrin (heme) portion of hemoglobin, (5) the hippuric acid and similar compounds found in the urms after ingestion of benzoic acid and other unoxidizable substances, and (6) the methyl group of choline, methionine, etc. Other aspects of glycine metaholism undoubtedly remain to he recognized.

Glycine is readily synthesizable by both the rat and the adult human; it is thus "dispensable" from the diet of these species. It has been shown, however, that about 1 per cent of glycine is necessary in the diet of the chick to ensure adequate nutrition (Almquist). This is one of several known examples of species specificity with regard to dietary amino acid requirements. The rate of synthesis of glycine in man bas been estimated by Quick, on the basis of the rate of excretion of hippuric acid after henzoate administration, to he somewhat over 0.5 g. per bour. It has been shown by Rittenberg and Schoenheimer that synthesized glycine may he derived from sources within the body as well as from the diet. It has been demonstrated that the amino acid serine is a significant precursor of glycine, and there is some indication that glutamic acid may also give rise to glycine, possibly through the intermediary formation of serine.

Knowledge concerning the metabolic degradation of glycine is ex-

tensive only a small portion will be reviewed litre. In the diabetic dog the carbon chain of glycine can be shown to be readily convertible into glucose. Animal tissues contain a flavoprotein enzyme which cataly zes the ovidation of glycine to glycyche acid and immonia

This reaction may be of importance in connection with the possible for mation of labile methyl groups by way of formaldehyde, as postulate on p. 1025

Of major interest is the part played by Alycine along with arginine and a labile methyl group in the biological synthesis of creatine. The separate steps in the formation of creatine have been shown to be a follows.

Bloch and Schoenheimer⁴⁹ demonstrated by feeding experiments using both glycine and arguine containing isotopic introgen that the two nitro-

gen atoms in the amidine portion (—(—)H) of creatine were derived from the amidine portion of arginine and the third introgen atom came from glycine Borsook and Dubnoff²⁰ showed that kidney tissue appears to form guanidoacetic acid from arginine and glycine and that his et usue probably synthesizes creatine from guanidoacetic acid in the presence of methionine Guanidoacetic acid itself the intermediate in creatine synthesis is found normally in only small amounts in animal tissues but readily gives rise to extra creatine formation when fed Glycine appears also to be directly concerned in the synthesis of glutathione the tripeptide which is thought to be of importance in intracellular oxidation and reduction it has been shown that when isotopic glycine is fed to animal the labeled glycine appears so rapidly in the glutathione of the animal body as to indicate the direct participation of glycine in the synthesis of his compound Involvement of glycine in the synthesis of heading the fine the synthesis of the compound involvement of glycine in the synthesis of heading the first properties.

^{**} Block and Schoenle mer J Bul (Iem 138 107 (1941)

18 B sook and Dut noff J B ol (hen 132 209 (1940) 138 389 (1941)

After prolonged administration of labeled glycine to an individual, the hemin isolated from the red blood cells contains sufficient isotopic carbon to indicate a major role of glycine in the synthesis of the pyrrole ring of the hemoglobin molecule. In fact the formation of heme from glycine has been found by in vitro studies in which glycine labeled with N¹⁵ and C¹⁴ was incubated with red blood cells of sickle-cell anemic patients or with the nucleated red cells of birds. Acctie acid also has been implicated in this synthesis (see p. 1009).

Although it was formerly believed that the animal was nuable to synthesize methyl groups and thus was forced to obtain preformed methyl groups for creatine, etc. from methionine or choline, it has now been demonstrated that many compounds are capable of supplying this metabolic essential. Thus choline, betaine, and methionine are able to coutribute labile methyl groups²¹ to each other and to guanidoacetic acid by the process of transmethylation, while glycine, seriue, methylol, formic acid, formaldehyde, and others are sources of methyl groups by conversion to an unknown one-carbon intermediate.

Relatively little work has been done on the role of glycine in the synthesis of the glycocholic acid of the liver. The relationship of glycine to hippuric acid formation is discussed in detail in Chapter 20. The ability of certain tissues to synthesize hippuric acid in vitro has been used as a

means of establishing possible precursors of glycine.

Alanine. This 3-carbon amino acid is readily synthesized by the animal body and hence is dispensable from the diet. Metabolism and synthesis are usually considered to involve the reversible equilibrium of oxidative deamination (see p. 1018) and transamination (see p. 1034). The reaction of oxidative deamination is as follows:

There is little evidence that this reaction actually occurs in the tissues; the L-amino acid oxidase of Blanchard, et al., attacks alanine but slowly, and there is no biological evidence for the synthesis of alanine from pyruvic acid and free ammonia. A more plausible metabolic pathway is by the reactions of transamination described on p. 1034, whereby synthesis would entail the transfer of the amino group of either glutamic or aspartic acid to pyruvie acid, to yield alanine and citber ketoglutaric acid or oxalacetic acid, and metabolic breakdown would be the reverse of these reactions. The pyruvie acid thus formed from alanine by transamination may then follow the normal course of oxidation of carbohydrate intermediates (p. 990). Conversion to pyruvie acid presumably is the explanation for the glucogenic action of alanine. By the use of alanine

⁷¹ See p. 1029.

¹⁸ Ulanchard, Green, Nocato, and Batner; J. Biol. Chem., 161, 583 (1945).

labeled with isotopic hydrogen (deuterium), it has been shown that alanine may be a source of acetyl groups in the acetylation of certain foreign amino acids exercted in the urine as acetyl derivatives (see p 1012 and Chapter 20).

Serine This hydroxy amino acid is readily synthesizable by the animal body and is thus dispensable from the diet. It nevertheless participates in a somewhat unusual variety of metabolic reactions in the tissues. In addition to its presence in proteins, it has been found in phospholipide material where it apparently serves in a manner analogous to choline and ethanolamine in the structure of phospholipide molecules (see Chapter 11). In fact, studies using serine labeled with isotopic nitrogen indicate that ethanolamine arises by the decarboxylation of serine, ethanolamic can then give use to choline by methylation. Serine is also a precursor of givenic (see p. 1025).

Serine is also concerned in the formation of cystine (see p. 1031). Serine is glucogenic in the diabetic dog. The "unnatural" optical isomer of serine, b-serine, is toxic to rats. The nitrogen of serine has been shown by isotopie experiments to be incorporated into the \gamma_r of urice and while the \beta-carbon appeared at C₂ and C₃ having been initially converted to formatic.

The anaerobic deamination of serine appears to take the following

$$\begin{array}{c|cccc} CH_1OH & CH_2 & CH_3 & CH_4 \\ \hline \downarrow & -H_2O & CH_2 & CH_3 & CH_4 \\ COH & COH_2 & C & CH_3 & CH_4 \\ \hline COOH & COOH & COOH & COOH \\ Serine & Amino & Imino & Pyruve acid \\ acid & acid & acid \\ \end{array}$$

Threonine This amino acid cannot be synthesized by the animal body, and its presence in the diet in adequate amount is required both for growth in the young rat and for the maintenance of nitrogen equilibrium in the adult human Threonine was the last 'essential' amino acid to be discovered in the sense that its isolation from casein and characterization by McCoy Meyer and Rose in 1935 first permitted the use of a mixture of pure amino acids as the sole source of introgen in an experimental diet.

Relatively little is known concerning the metabolism of threonine It is glucogenic and antiketogenic. It is not attacked by the L-amino aco oxidase of rat kidney, but there may be another enzyme concerned since it has been reported to be dearminated by kidney this under anaerobic conditions to produce α ketobutyric acid CH_1 CH_2 COOOH Only the optically active form found naturally is utilizable for growth Threome and by since are the only amino acids known which do not transaminate.

Methionine This sulfur-containing amino acid cannot be synthesized by the animal body from the ordinary constituents of the diet and mutbe present in adequate amount for the promotion of growth in the young

animal and for the maintenance of nitrogen equilibrium in the adult human. Methionine alone will satisfy all of the sulfur requirements of the animal, since it is readily converted into cystine in the tissues, as described on p. 1031.

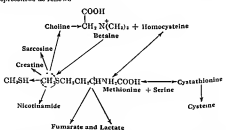
Methionine is glucogonic in the diabetic dog. Metaholism may proceed in part by deamination through the corresponding keto acid, CH₂·S·CH₂·CC+·COOH, since this latter compound is formed to considerable extent on incubating methionine with liver tissue, and the young rat will grow on a diet containing the keto acid in place of methionine. Further stages in the oxidative breakdown of methionine are obscure. On complete oxidation, the sulfur is found in the urine as sulfate, either inorganic or ester, as with cystine (p. 1032).

TRANSMETHYLATION. An important metabolic function of methionine is in connection with the process known as transmethylation. It has been shown by du Vigneaud and associates that the animal body is able to use the methyl group of methionine to methylate certain nitrogenand sulfur-containing compounds of the body. Methionine, choline, and betaine (trimethylglyciue) have been shown to be suitable dietary sources of such methyl groups; of these, methionine appears to be quantitatively the most important. In the biological synthesis of creatine, for example (see p. 1026) the methyl group of creatine is derived by transfer from methionine. The process of methyl transfer from one compound to another is called transmethylation. A methyl group which may be so transferred is called a labile methyl group; thus far, only methyl groups attached to either nitrogen or sulfur appear to be labile and to participate in transmethylatiou. Contrary to the former opiniou, labile methyl groups need not be supplied in the diet but can be synthesized from glycine, serine, formic acid, and their precursors (see p. 1025).

The role of methionine in transmethylation has been adequately proved by the use of methionine synthesized to coutain isotopic hydrogen (dcuterium) in the methyl group. When such labeled methionine is iucluded in the diet of an animal, the methylated compounds choline and creatine subsequently isolated from the animal tissues prove to contain sufficient isotope to justify the conclusion that methyl groups have been transferred from the methionine to the other compounds in question. The ability of methionine to furnish methyl groups for the synthesis of choline from suitable precursors explains the lipotropic effect of this amino acid (see p. 1012). The transfer of methyl groups from methionine to choline is reversible, most probably through the intermediate formation of the amino acid homocysteine (see below); methylation of guanidoacetic acid to form creatine is irreversible, and the constant excretion of body creatine as nrinary creatinine represents a loss of methyl groups from the body. The daily requirement for methyl groups, however, appears to be considerably in excess of such urinary loss, so that other pathways of methyl group degradation must be present; relatively little is known about this at the present time. Most of the evidence for transmethylation has been obtained with the rat, but the process has also been shown to occur in the adult human.

After removal of the methyl group from methionine, the amino acid

homocysteme results Homocysteme, HS CH2 CH2 CH(NH2) COOH and its disulfide form homocystine (analogous to the relation between cysteme and cystine) are synthetic amino acids which have not as yet been isolated from natural sources. Their biological availability is such, however, as to lead to the behelf that they represent normal intermediates in methionine metabolism If an animal is placed on a diet containing no methnomine but adequate amounts of homocysteine or homocystine, together with a source of labile methyl groups such as choline, betaine, or serue, the animal will synthesize the methionine it needs for normal growth If the methyl groups of the dietary choline are labeled with deuterium, the methionine subsequently isolated from the tissue proteins proves to contain the isotope in its methyl group. The animal body 15 therefore capable of transferring methyl groups from choline via betaine to homocysteine, to form methionine, as has already been pointed out, this transfer is reversible Since the reversible exchange of methyl groups between methionine and choline eau be demonstrated not only on the homocysteme diet but also when the diet contains adequate amounts of methionine and choline, it is believed to represent a normal metabolic process of the tissues The implication of homocysteine in the formation of cystine from methionine is discussed on p 1031. The general metabolic relation between methionine, homocysteine, and related compounds may be represented as follows



In addition to its known metabolic functions methionine appears to play a specific part in protecting the byer from damage by such poisons as carbon tetrachloride phosphorus, arsenic, and chloroform. The mode of action here is unknown

Cystine and Cysteine. These two sulfur-containing amino acids are usually considered to be metaboheally equivalent since one may be so readily convicted into the other by oxidation or reduction. There is some cysteine and cystine particularly in specific organs such as the liver and kidney. In the rare metabolic ahnormality known as cystinura, individuals regularly excrete significant amounts of cystine in the urine. This

exerction continues during fasting, and is not increased by the oral administration of cystine itself; the feeding of cysteine or of methionine does, however, augment cystine exerction. Thus the cystinuric distinguishes between orally administered cystine and cysteine, and the normal individual may do so in specific tissues as well.

It was thought for a long time that cystine was an indispensable amino acid, but Jackson and Block showed in 1932 that methionine could replace cystine for growth purposes in the rat on a low-cystine diet, and the ability of the animal body to form cystine from methionine is now well established. In 1939 Tarver and Schmidt demonstrated, by the use of methionine containing radioactive sulfur, that the sulfur of cystine was derived from the sulfur of methionine

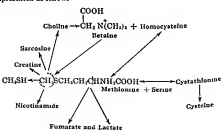
The mechanism of formation of cystine from methionine has heen the subject of considerable study, and appears to have been finally established by the work of du Vigneaud and his associates as follows; methionine is demethylated to form the amino acid homocysteine; this condenses with scrine to form an unsymmetrical thio-ether, called cystathionine; the cystathionine undergoes enzymatic cleavage so that the sulfur remains with the serine moicty, to produce cysteine. The fate of the remaining portion of the cystathionine molecule is as yet not known. These various steps may be illustrated as follows:

Evidence for the conversion of methionine to homocysteine has already been presented (p. 1029). When homocysteine and scrine are inenhated with liver tissue, cysteine is formed. It has also been shown by Stetten that when serine labeled with isotopic nitrogen is fed to animals, the cystine isolated from the tissues contains such a high proportion of the isotope as to indicate (but not prove) the direct conversion of serine to cysteine. Cystathionine, the postulated intermediate, has been found in the livers of fasting rats which were fed both L-methionine and L-serine. If either amino acid were fed alone, no cystathionine was observed. If methionine is synthesized to contain both isotopic sulfur and isotopic carbon, and this doubly labeled compound is fed to rats, the cystine isotopic sulfur, but no isotopic earbon. This is further evidence that it is

⁷⁴ Hess: Arch. Brochem Braphys , 40, 127 (1952).

¹⁴ du Vigneaud, Kiliner, Machele, and Cohn. J Biol. Chem., 155, 645 (1944).

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Hess Arch Buchen Biophys 40 1. (1952)
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the sulfur and not the carbon chain of methionine which is involved in the formation of cycline

Cystine and cystenic contribute to the formation of many important sulfur containing compounds in the immal body. Cystenic is a component of plutathione (y plutamyleyster) also may appears to be the precursor of the trumne of the taurocholic icid of the bile probably through intermediate oxidation to form cyster acid.

Cysteme is likewise found in the urine in combination with certain unoudizable substances (detoxication) to form what are called mercapture aculs (see Chapter 20). Cysteme is glucogenic in the animal body possible through the intermediate formation of senne. On complete oxidation of cysteme and cystine the sulfur is found in the urine either as morgane sulfate or to some extent as organie esters of sulfure acid with such compounds as indoxyl, phenol, etc. This is also diccussed in Chapter 20.

ONE-CARBON INTERMEDIATES In the discussion of the metabolism of glyenic serine methonine cystine and histidine mention is made of labile methyl groups and one-carbon intermediates. Whether these are

CH.

CH₂

the same compounds or not, it is known that they are metabolically related and for purposes of simplification have been formulated as HCOOH in the diagram on p 1032 which illustrates some of the metabolic pathways of these oue-curbon intermediates

Leucine and Isoleucine Relatively little is known concerning the metabolism of these two amino acids. They are not synthesizable from dietary constituents and are required in the diet for the growth of young animals and the maintenance of introgen equilibrium in the adult Leucine and isoleucine are ketogenic rather than glucogenic runnio acids. The conversion of leucine to acctoacetic and possibly proceeds as follows.

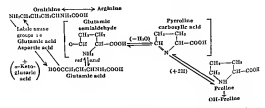
Value. This amino acid is required in the diet in adequate amount, as evidenced both by growth studies with young animals and by the maintenance of introgen equilibrium in the adult human \(\Lambda\) possible metabolic pathway involves deminiation to form \(\alpha\) ketosovaleric acid, since this compound will replace value in an experimental diet

CH₂

Value is convertible, in part at least, to glucose in the disbetic dog

Proline and Hydroxyproline. These two amino acids can be synthesized by the animal body from dietary or tissue precursors, and proline is glucogenic in the dibetre dog: A probable precursor of proline is or inthine, since ormitline and proline are interconvertible in the organism proline metabolism may proceed through oriuthine formation. Glutamic acid has likewise been shown to arise from proline, further metabolism via this pathway would then be that of glutamic acid, and this may be the origin of the glucogenic action of proline. Hydroxyproline can be synthesized from dietary proline, there is some evidence that this reaction is not receivable, and that the further metabolism of hydroxyproline mandatory.

not be similar to that of proline. These various relationships may be summarized as follows.



Glutamic Acid. This dicarboxylic amino acid is present in fairly large amount in many animal and vegetable proteins. It is readily synthesizable by the animal body, and is thus dispensable from the dict, it is probable, however, that the case of synthesis within the tissues is more properly a reflection of the important part played by glutamic acid in the general processes of nitrogen metabolism within the cell rather than an indication of a "nonessential" function

Evidence that glutamic acid may be an important intermediate in general nitrogen metabolism is afforded by the work of Schoenheimer and associates using various amino acids labeled with isotopic nitrogen (Nⁱⁱ). When such labeled amino acids are fed to an animal, the isotopic nitrogen is found not only in the fed amino acid incorporated into the tissue proteins of the animal, but also to a significant extent in various other amino acids as well, and of these, glutamic acid usually exhibits the highest concentration of isotope it is probable that transamination (see bellow) is partly responsible for this, the presence of glutamic acid in glutathors may also be connected with the apparently rapid transfer of detary amino nitrogen via glutamic acid.

Transamination An important metabolic aspect of glutamic acid is its participation in the reactions of transamination. In the transamination reaction the amino group of glutamic acid is transferable to either private or oxalacetic acid to produce a-ketoglutanic acid and the amino acid alamine or aspartic acid, as the case may be These reactions are reversible so that glutamic acid may be synthesized from a-ketoglutaric acid and either alamine or aspartic acid. The diagram on p. 1035 illustrates the transamination reactions.

Enzymes catalyzing the upper reaction (glutamic aspartic transaminase) and the lower reaction (glutamic alanne transaminase) have beer isolated from animal tissues, and are believed to contain a pyridounic (vitamii B) derivative as prosthetic group. A third postulated reaction between aspartic acid and pyriuve acid to produce ovalacetic acid and alanine, is due, according to Green, to the presence of both of the transaminases mentioned. It is noteworthy that no free animonia is formed during transamination, the reaction apparently involving condensation of

amino acid and keto acid through the amino and keto groups to form an intermediate which is then split in such a way that the amino nitrogen is transferred from the original amino acid to the keto acid chain.

In addition to transamination, glutamic acid likewise undergoes reversible oxidative deamination.

Glutamic acid ≈ α-Ketoglutaric acid + NH;

The reversal of this reaction is one of the few known examples of the biological synthesis of an amino acid from the keto acid and ammonia, and may represent one of the pathways for the demonstrated (although limited) couversion of dictary ammonia nitrogen into amino acid nitrogen.

As would be expected from the equilibrium between glutamic acid and its keto acid, a-ketoglutaric acid, which is an intermediate in earho-hydrate metabolism (p. 990), glutamic acid is glucogenie in the diabetic dog. Certain other amino acids such as proline, ornithine, and histidine are known to give rise to glutamic acid in metabolism, and this may also explain their glucogenie action. The other heto acids involved in transamination (oxalacetic acid and pyruvic acid) are likewise carbohydrate breakdown products, and this fact may represent an important link between amino acid metabolism and carbohydrate metabolism.

Glutamic acid is combined as one of the three portions of the molecule of the vitamin pteroylglutamic (folic) acid, the other two being p-aminobenzoic acid and pteroic acid; the various conjugated forms of this vitamiu are due to the multiplicity of glutamic acid units present in peptide linkage—e.g., the conjugate in liver contains seven such units while that obtained from fermentation contains three. (See Chapter 35.)

Glutamic acid appears to he present in proteins largely in the form of its amide, glutamine, IIOOC-CII(NI₂)-CH₂-CH₂-CONI₂. Free glutamine has been found in small amount in the hlood, and according to Van Slyke and associates, glutamine or a glutaminelike compound is of importance as a precursor of urinary ammonia. Glutamine has also been implicated

in the process of urea formation by the liver but its role here, if any, is obscure the reported presence in cancer tissue of the "unnatural optical isomer of glutamic acid, p-glutamic acid is found in the capsule of the anthrax bacillus and in other products of bacterial origin.

Glutimic acid is decarboxylated in the α position to give γ aminobutine acid Glutamic decarboxylases are widespread in plants and bacteria, but have been found only in the higher portions of the central nervous system of animals. The importance of glutamic acid, glutamice and γ aminobuty no read in the metabolism of the brain is not yet clear but the present evidence indicates that these compounds must play a basic part in the physiology of nervous tissue

Aspartic Acid The metabolic aspects of this amino acid are less well understood than those of glutamic acid but it appears to be similar to the latter in certain respects. It is readily synthesizable within the body and shares with glutamic acid in the transfer of dictary amino nitrogen as described on p. 1034, but to a lesser extent. Synthesis probably proceeds via the transamination reaction involving the keto acid oxalacetic acid and glutamic acid as described previously. Metabolic breakdown by the reversal of this reaction to form oxalacetic acid, which is a carbobydrate intermediate serves to explain the glucogenic power of aspartic acid as well as to indicate the pathway of further degradation. The amide of aspartic acid aspartic acid aspartic acid asparagine. HOOC CII(NH2) CII; CONH2, is found in tissue proteins but its significance is obscure.

Histidine Histidine is required in the diet of the young growing animal but is apparently dispensable from the diet of the adult himan as established by the maintenance of introgen equilibrium in man on diets free of histidine Here as with argume it may be that rate of synthesis is the limiting factor with the possibility that bacterial action in the intestinal tract may also be concerned. Histidine is apparently metabolized via the pathway shown at top of p. 1037.

In addition to its presence in tissue proteins histidine is found in the animal body in the muscle constituents earnosine (β -alanyl histidine) and anserine (β -alanyl michlyl histidine) and it is probably a precursor of the red blood cell constituent ergothioneine which is a betaine of thiolistic dine. Histidine is regularly found in the turne during preemancy (and also

[&]quot; Edlbacker and Kraus Z physiol Chem. 191 225 (1930)

in some other conditions); the significance of this is unknown. Histidine is glucogenic, possibly because it gives rise to glutamic acid during metabolism.

Decarboxylation of histidine produces histamine:

Histamine is an organic base with a very powerful pharmacological action on certain body structures. In addition to its well-known effect on blood pressure (capillary dilatation), it is a potent stimulant of gastric secretion (see p. 360), and has been implicated in the complex symptomatology of anaphylactic shock and in eclampsia. Drugs which have an action specifically antagonistic to that of histamine are known as antibistamines; many if not all of those compounds are structural analogs of histamine and presumably owe their effect to an autimetaholite action (see Chapter 36).

Lysine. This amino acid is required in the diet of the young growing animal and for the maintenance of nitrogen equilibrium in the adult human. It is not glucogenio in the dabetie dog. It is unique among all the amino acids thus far investigated in that it does not appear to be capable of obtaining its nitrogen from other dietary sources, as can be done by certam other amino acids (see discussion on p. 1022). Lysinc, however, can contribute its nitrogen to other amino acids after metabolic breakdown.

The metabolism, therefore, may involve an oxiditive dealing it on which for some reason is irreversible. This is borne out by the fact that neither the keto and nor the hydroxy and deny time of lysine our substitute for this amino and in experimental diets. Studies have indicated that lysine may first form a aminoadipie and which is then converted into glutanized and hence to glutanic real.

Arginine. The question of whether or not this amino acid can be synthistized by the animal body has been the subject of considerable debate in the past it now appears to be settled largely through the work of Rose, using diets containing mixtures of pure amino acids that arginine can be synthesized by the tissues of the voing growing rat for example but not at a rate fast enough to supply the needs of the animal for optimal growth it is thus indispensable in this sense in the diet of the voing growing rat. The adult rate (and the adult human) apparently can supply arginine by synthesis from other sources at a rate adequate for nutritional needs. On the other hand, the requirements of the click for arginine are such that this amino acid must be considered indispensable in the definition of the control of species differences with respect to amino acid requirements.

Various studies using amino acids labeled with isotopic nitrogen and hydrogen have indicated that arginine is synthesized in the body by the

addition of an amidine group (—C=NI) to ornithme HOOC CH(NII)—CH₂ CH₂ CH₂ XII₂ The origin of the amidine group is not known (a possible mechanism is by the ornithme cycle discussed below), but ornithme is readily synthesizable biologically—for example from rolande (1033). Evidence that the synthesis of arginine is reversible is not available, but it is known that the amidine portion may be transferred to glycine to form guanidoacetic acid in the synthesis of creatine (p. 1026). Arginine also may be degraded to form ornithme and ures in the presence of the enzyme arginase as described below. There is evidence that arginine may give rise via ornithme to proline hydroxyproline and glutamic acid, and these may represent further stages in metabolism. Both arginine and ornithme are glucogenic, possibly because of their convertibility to the amino acids just mentioned. There is some evidence that the α amino.

nitrogeu of arginine does not undergo oxidative deamination to form the corresponding keto acid, or if so, that this process is irreversible.

Interest in the metabolic function of arginine has been directed primarily toward its possible relation to the synthesis of creatine and the formation of urea. The role of arginine in creatine formation is described in detail on p. 1026. Interestingly enough, an arginine analog of phosphocreatine (namely, phosphoarginine) appears to serve in place of phosphocreatine in the muscles of certain lower animal species.

The possible role of arginine in urea formation hy mammalian liver has been evident for a long time heeause of the presence of the enzyme arginase in liver; arginase catalyzes the hydrolytic splitting of arginine into ornithine and urea. According to Krehs and Henseleit, arginine enters into a cyclic mechanism, along with the amino acids ornithine and citrulline, for the conversion of ammonia into urea in the liver. On the hasis of later developments, the process is now pictured as follows:

According to this view, ammonia and carbon dioxide are added to the amino acid ornithine to produce the amino acid citrulline. On the further addition of ammonia to citrulline, arginine is formed which then undergoes action by the enzyme arginase to produce urea and to regenerate ornithine, which can then proceed through the cycle again. The original evidence for this theory was based largely on the demonstration that both ornithme and citrulline catalyzed the synthesis of urea from free ammonia hy liver slices. Additional evidence tending to support the theory has come from several sources. If the liver slices are incubated in the presence of bicarbonate containing isotopie earbon, the earbon isotope is found in the urea synthesized. Schoenheimer's work on the distribution of isotopic dictary nitrogen between the amidine moiety and the ornithine moiety of arginine isolated from tissue proteins, and the isotope content of the urinary urea, led him to conclude that the results supported the ornithine cycle theory. According to Gornall and Hunter, the accumulation of citrulline in liver tissue during urea formation under special conditions may be demonstrated, and citrulline may also be found in small amount in the blood. Beadle and Tatum also have presented evidence

that the ornithme-cycle mechanism functions in the red bread m

Tryptophan. This amino acid is required in the diet for growth the young animal and the maintenance of nitrogen equilibrium in adult human It has several metabolic pathways which appear to v. somewhat in different species of animals. One of the most interest findings in recent years is that tryptophan is converted in the body to vitamin, nicotinic acid (niacm). Other end products appear to be alam anthranilic acid, kynurenic acid, etc. (cf. diagram). The quantitat aspects of these known metabolic pathways have not as yet been clidated, but it appears that the conversion to nicotinic acid plays a m role, for it requires from 50 to 300 g. of tryptophan to make 1 g. of vitamin.

Kynurenic acid Xanthurenic acid
Tryptophan Kynurenine | Raboffavia | 3-Hlydrozykynurenine | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenisec²⁴ | Kynurenise

urine after the ingestion of large doses of tryptophan (Jackson). There is some basis for the belief that kynurenic acid is au eud product and not an intermediate in tryptophan metaholism. Kynurenine is claimed to be a hormone for Drosophila. Tryptophan is converted into 4,8-di-bydroxyquinoliue-2-carbo\ylic acid (xanthurenic acid) by the rat on a diet deficient in vitamin B₆. The red bread mold Neurospora causynthesize tryptophan from indole and serine. The reverse of this reaction, i.e., the production of indole by bacterial action ou tryptophan in the intestinal tract, is believed to be the origin of the indole, skatole (methyl indole) and indican of the feces and urine. Tryptophan deficiency produces a type of cataract in the rat, and certain evidence indicates a relation between anemia, tryptophan, and vitamin B₆ (pyridoxine).

Tryptophau cataliolism may also proceed by oxidative deamination to the corresponding keto acid, indolepyravie acid, since this compound (and indolelactic acid) will replace tryptophan in experimental diets. Tryptophan is apparently not ketogenic and will alleviate experimental ketonuria

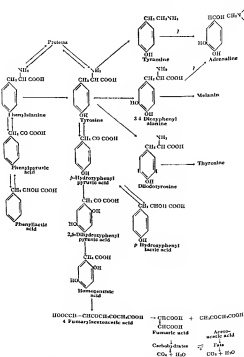
in the rat.

Phenylalanine and Tyrosine. These two aromatic amino acids usually are considered together because of their close metabolic relationship. Of the two, phenylalanine cannot be synthesized by the animal body (except from its corresponding keto-or hydroxy-acid, which are not ordinary constituents of the diet) and therefore must be present in the diet in adequate amounts. The needs of the animal for tyrosine, however, can be readily supplied by dietary phenylalanine, as established by growth studies with young rats and by the maintenauce of nitrogen equilibrium in the adult human. Phenylalanine, therefore, is convertible into tyrosine by the organism; this conversion has been directly demonstrated by the feeding of phenylalanine labeled with isotopic hydrogen (dcuterium) to rats, followed by the subsequent isolation of tyrosine from the animal's tissue proteins; such isolated tyrosine contained sufficient deuterium to establish its origin from the fed phenylalanine. All available experimental evidence indicates, however, that the conversion of phenylalanine into tyrosine is irreversible; tyrosine cannot be converted into phenylalanine in the animal body.

Knowledge concerning the intermediary metabolism of both phenylalanine and tyrosine is still far from complete. In addition to their roles as essential components of the body proteins, these two amino acids, and particularly tyrosine, appear to be concerned with the formation of a variety of physiologically important substances such as thyroxine, adrenaline, the pigment melanin, and others. Certain of the known or postulated metabolic interrelationships of tyrosine and phenylalanine are sum-

marized in the diagram on p. 1042.

Both phenylalanine and tyrosine are ketogenic amino acids, i.e., they can be shown to give rise to acctoacetic acid under the proper conditions, and neither amino acid is glucogenic. It is postulated, therefore, that after the loss of nitrogen which is evereted as urea, both phenylalanine and tyrosine are normally oxidized to carbon dioxide and water through the intermediate formation of acetoacetic acid. Whether these two amino acids follow a common metabolic pathway (i.e., phenylalanine -> tyrosine



→ intermediates → CO₂ + H₂O) is a matter of dispute Possible direct metabolities of phenylalanine other than tyrosine include phenylpyruvic and phenyllactic acids phenylpyruvic acid is found in the urine of animals after feeding with phenylalanine

I olling reported that a certain proportion of individuals with mental disease regularly exercte phenylpyruvic and in the urine This condition has been called oligophrenia phenylpyruvica. If phenylalaniue is administered to individuals suffering from this metaholic abnormality, the excretion of phenylpyruvic acid is inereased. Thus there is good evidence that phenylpyruvic and phenyllacticacids are produced from phenylalanine; there is equally good evidence that the reactions leading to their production are reversible; for instance, both phenylpyruvic acid and phenyllactic acid can replace phenylalanine in the diet of the growing rat.

The metaholic fate of tyrosine is somewhat hetter understood than that of phenylalanine; the irreversible blockage between tyrosine and phenylalamne permits a more precise estimation of the significance of possible intermediates obtained under experimental and ahnormal conditious. As with phenylalanine, possible metabolites of tyrosine include the corresponding keto- and hydroxy-acids, p-hydroxyphenylpyruvic and p-hydroxyphenyllactic acids, respectively. Both of these compounds have been isolated from human and animal urine under certain conditions. Medes has described a condition called turosinosis (only one such case has thus far heen found) in which an adult human excreted significant amounts of p-hydroxyphenylpyruvic acid in the urine. The output of this compound was augmented with increased intake of protein or of pure tyrosine or phenylalanine; feeding the compound itself resulted in the unchanged excretion of most of it Leviac and associates discovered that the infant human may exhibit a spontaneous defect in tyrosine metabolism characterized by the excretion of significant amounts of p-hydroxyphenylpyruvic and p-hydroxyphenyllactic acids, in the presence of a low vitamin-C intake and an excessive intake of tyrosine or phenylalanine, either in the form of protein or as the pure amino acids Vitamin-C administration aholished the defect except in the presence of excessive amounts of phenylalanine or tyrosiae, Sealock and associates have shown that the scorbutic guinea pig excretes tyrosine metabolites (p-hydroxyphenylpyruvic acid, homogentisic acid) after administration of tyrosine or phenylalanine; vitamin-C administration prevented the exerction of these compounds. It has also been shown that scorbutie guinea-pig liver is unable to oxidize tyrosine in vitro except in the presence of vitamin C.

It would appear from these and other findings that the keto acid of tyrosine is a normal intermediate in tyrosine metabolism, and that vitamin C is concerned in the further utilization of the keto acid. The possibility remains open, however, that oxidative deamination of tyrosine is only one metabolic pathway and that the metabolism may follow some as yet unknown sequence of reactions.

Further knowledge concerning tyrosine metabolism has been afforded by the study of the metabolic abnormality known as all.aplonuria. In this relatively rare condition, individuals exerte the compound homo-

[&]quot;This disease is characterized by the fact that the individual has lost the ability to convert plony) alianne into tyroune. It is consequence phenylalanue must be omitized via the pathway phurislalanue — phenylaprune and — phenylaprune and — phenylaprune and phenylaprune and phenylaprune and a neurotoxic agent which is detoxified by combination with glutanine to yield phenylacety, lebutanine A pointed out on p. 1630, glutanine and and glutanine appear to be of special significance in the metabolism of the higher centers of the central nervous system. All patients with objective and phenylaprune are mentally defective. Is this due to a chronic deficiency of glutanine?

gentisie acid (for structure see diagram on p 1042) in tho unne 8 In the presence of alkalı and oxygen homogentisic acid forms a dark brown or black pigment and it is the darkening of the urine under these conditions which usually leads to the discovery of alkaptonuria Homogentisic acid is believed to be a normal intermediate in tyrosine metabolism probably through the intermediary formation of 2 o-dihydroxyphenylpyruvic acid (see diagram p 1042) and the metabolic defect in alkaptonuma is thought to be the lack of ability to carry the oxidation beyond the stage of homogentisie acid formation Feeding of tyrosine or phenylalanine to an alkaptonurie increases the output of homogentisic acid and fed homogentisie acid is exercted unchanged. The normal individual is able to oxidize homogentisic acid completely Alkaptonuria may be produced experimentally in animals (and in man) by prolonged or excessive feeding of either tyrosine or phenylalamine and in scurvy Vitamin C administration will correct the alkaptonuria of the scorbutic guinea pig but has no apparent effect on human alkaptonuma

Certain other aspects of tyrosine metabolism are of physiological importance. Both the diodotyrosine and thyroxine of the thyroid gland are considered to arise from tyrosine. Both of these substances have been isolated from casein and other tyrosine-containing proteins treated in surro with alkaline iodide solution and the incubation of thyroid gland slices in the presence of radioactive noidide as a tracer results in the production of diodotyrosine and thyroxine containing the radioactive material. Epinephrine is likewise thought to originate from tyrosine it has been shown that animal tissue can decarboxylate tyrosine to yield tyramine oxidation and methylation of this compound could produce epinephrine? In alternate possible pathway would involve the intermediate formation of 3+chhydroxyphenylatanine followed by decar boxylation to yield hydroxytyramine and oxidation and methylation to produce eninephrine.

According to Bloch and to Raper it would appear that the pigment melanin is related to tyrosino metabolism. Melanin is not a simple chemical substance but rather a mixture of pigments of ill-defined composition. It is a normal skin pigment and appears to be low or lacking in the condition known as albinism. It is produced in excessive amounts by melanote tumor cells and may even be excreted in the urine (melanitra) under these conditions. It is also produced in excessive amount in Addison side case (bronzed diabetes). Bloch showed some years ago that the skin contains an enzyme (dopase) capable of converting the compound 3+dhydroxyphenyilalanine (dopa) into melaninlike material and that this enzyme was absent from the skin of albinos. According to Raper the probable precursor of 3-d-dihydroxyphenyilalanine is tyrosine.

and discard the urine Prepare a thoroughly clean bottle of proper size, introduce into it sufficient toluene to cover the bottom of the hottle, and use this bottle for the collection of all urine voided during the following 24 hours, including the urine obtained by emptying the bladder at the close of the 24-hour period, e.g., at 8 A V the next day During the day, when not actually in use for the introduction of a urine fraction, the hottle should be kept in a refrigerator or cold room in order that the sample may not ileteriorate before it is examined. Measure the volume of the sample and determine its specific gravity (see p. 785) and reaction before proceeding to the quantitative estimation of any specific urinary constituents.

For metabolism work on dops, a eage is used which is provided with a screen hottom. Below this is a tray which slopes toward a central hole through which the urine passes into a bottle containing toliene Each day's output is collected and filtered to remove hair, etc., and may then be diluted to a definite volume, usually 500 or 1000 ml. This procedure facilitates subsequent calculation.

To obtain the complete 24-hour secretion of urine of dogs, catheterization must be resorted to Because of the difficulty of eatheterizing male dogs, bitches (especially those who have had pups, and hence have stretched vaginas) are used for this type of experimentation. Care must be taken to have the eatheter sterile, in order to avoid infection or cystitus Rubber eatheters are sterilized by holing in water, while metal oglass catheters may be washed and kept in alcohol. The simplest procedure is to use a speculum (a nasal dilator is about the right size) to stretch the vagina had then to insert the eatheter directly into the urethra, avoiding contact with the vaginal wall. The exact location of the urcthral orifice can be determined with a little experience. Force should never he used. After the bladder has been complied, it is washed out several times with warm water, the washings heing collected with the urine and made up to volume as described above. Finally, if it does not interfere with the experiment, it is desirable to introduce about 50 ml of warm saturated hore need solution into the hladder from time to time.

The most satisfactory method for obtaining urine from n rabbit is to hold its head up hetween one's laces and to npply gentle pressure on the lower abdomea. By this process of "milking," practically all the urine may be obtained

For the collection of urine in metabolism work on rais, a metabolism cage (see Appendix) is employed

2. Complete Analysis of Urine. Ingest an ordinary mixed diet (or any special diet) and collect the urine accurately for a 24-hour period (see above). Measure the volume of the sample, determine the specific gravity, and preserve the urine (see above) until the following constituents have been determined (for methods of analysis, see Chapter 31), total solids, titratable acidity, hydrogen-ion concentration, total nitrogen, amino acid nitrogen, ammonia, urea, uric acid, creatinine, total sulfur, ethereal sulfates, inorganic sulfates, neutral sulfur (by difference), total phosphates and sodium chloride.

Calculate the nitrogen and sulfur partitions, i.e., the percentages of the total nitrogen and sulfur which occur in the different forms, and tabulate the data from the complete analysis. Compare your results with those listed in the tables on pp. 1955 and 1657

sired to economize time and effort in the chemical examination, the daily fecal output or an aliquot portion of each stool may be collected in a friction-top can or pail of suitable size and preserved by thyinol and refrigeration. This method has been found satisfactory when the feces are to be examined for morganic constituents or total introger. For the determination of fat, earbohydrate, etc., the fresh stool should be employed because of the possibility of error due to by drolysis of fat to fatty and, etc.

In the preservation of feces for the determination of total attrogen, the following simple procedure may be used. Introduce each stool into a weighed friction top can or pail and place the vessel in a cold room or refrigerator preferably, near or below 0° C. At the end of the period, max the feces thoroughly and analyze weighed portions In case individual stools are analyzed, the stool should be collected in a verified flat-bottomed porcelain dish. We there mixing the feces very thoroughly, the weight of dish, spatula, and feces is determined and the weight of the feces secured by difference. 1° A portion of the well mixed feces is then introduced into a large weighing bottle containing a glass how Desired amounts of feces are then removed for analysis and the exact weight of such amounts obtained by difference.

The daily output of feees in dogs is quite variable, so that in metabolism work it is advail to collect feees in princh, rather than each day. The stools are collected from the series floor of the cage and placed in weighed pass. They are fined by adding alcohol, stirring and exporating on the water hath, and then weighed. A few drops of sulfurne acid should be added to present the loss of ammonia. After drying the feees may be ground in a infli or mortar, to facilitate uniform sampling.

Experiment ingest a uniform diet for four days, or longer for greater accuracy. Divide the interval into two equal periods and separate the feces by charcoal or carmine (see Exp. 4 above). During the second period ingest 10 g of agar agar at each meal. Collect the feces for each period (see Exp. 4, above) and note the increase in the daily exerction under the influence of the agar ingestion. What was the increase per gram of agar?

II EXPERIMENTS ON NITROGEN METABOLISM

I Time Relotions of Protein Metabolism. It is a well known physiological fact that an interval clapses between the ingestion of protein food and the appearance in the urine of certain products representing the complete criabolism of this food. The chief among these is urea. The term nitrogen lag h is been used to designate the period clapsing between the ingestion of protein and the exerction in the urine of a quantity of nitrogen equivalent to that contained in the protein.

Experiment ingest a simple uniform diet whose exact composition has been determined by analysis or whose approximate composition has been estimated (See Appendix for table showing composition of foods) Continue this diet from i to 4 days Collect the urine in 2 hour periods from 7 A M to 11 P M and in an 8 hour period between 11 P M and 7 A M Analyze each specimen for total nutrogen or urea At the end of this preliminary period add to the uniform diet, at one meal, a weighed quantity (150 250 g) of lean meat specially prepared and analyzed Collect the urine in periods as before and determine total nitrogen or urea excretion, tabulate the data and plot curves showing the course of the nitrogen excretion on the various days of the experiment How long was the nitrogen and '?

A less accurate experiment than the above but one which yields interesting data may be carried out as follows

Ingest a simple diet whose nitrogen content can be estimated with some degree of accuracy Collect the urine in 2 hour periods from 7 a Mt to 11 Mt and in an 8 hour period from ii P Mt to 7 a Mt and analyze for total nitrogen or urea. The next day ingest the same diet plus 150-250 g of ienn meat whose nitrogen content has been determined by analysis or estimated. Collect the urine as upon the previous day and determine its total nitrogen or urea content. Plot curves showing the course of the nitrogen or urea excretion on each of the days. How soon after the ingestion of the large quantity of meat did you note an increase in the nitrogen or urea excretion? How many hours after the meal was the maximum quantity of introgen or urea excreted?

2 Digestibility and Biological Value of Protein The term protein utilization or biological value is used to indicate the percentage of the ingested food introgen actually assimilated 'disorbibility or digentibility indicates the percentage of the ingested food nitrogen which is absorbed.

The digestibility of food protein may be approximated by the following procedure Ingest any diet of known nitrogen content for a period of three days (see Appendix Composition of Foods) Longer periods are necessary for accurate work. Collect all feces from the diet, making the separations as directed on p. 1046 using carmine as the initial marker and charcoal as the float marker or vice versa. Preserve the feces as directed on p. 1045 Mix the total feces thoroughly and determine the nitrogen by the Kjeldahl method. The approximate nitrogen utilization may be calculated as follows.

Insamuch as the introgen content of the feces does not originate entirely from the food but represents in part residual metabolic products ie, intestinal epithelium, lactina secretions etc., a correction is usually made for metabolic nitrogen in more exact work (see p. 1020). The value thus obtained is more properly designated as the percentage digestibility or the coefficient of digestibility.

To determine the true percentage digestibility of food protein, proceed as follows Ingest a nonnitragenous diet as described on p 1055 for a period of two days, using sufficient agar agar to insure a daily fecal output which shall approximate in weight that obtained when the regular protein diet was in gested "Separate and preserve the feces as directed on p 1046 Mixthoroughly and analyze for nitrogen according to the Kieldahl method. This gives the metabolic fecal nitrogen Folium this period with a test period during which the protein in question is included in the diet Calculate the percentage digestibility of the prutein of the diet as follows

The estimation of protein utilization in the body must take into account not only absorptic if from the intestine, but also that fraction of the absorbed introgen which is retained.

Mitchell ** who has used this procedure extensively in the biological evaluation of proteins felt that this could be done by determining the urinary introgen exerction on a nonintrogenous dirt (indegenous introgen) and subtracting this from the urinary nitrogen on the protein-containing due to obtain the urinary introgen settally due the metabolism of the protein. It is not clear that this procedure is entirely said hased as it is upon Folin's cleasing distinction between endogenous and exogenous metabolism which now appears incorrect (see discussion on p. 1021) Nevertheless the procedure under the conditions presented by Mitchell is capable of giving reasonable and consistent results whatever may be its theoretical basis. The nitrogen balance method has been employed by Rose to determine the indispensable amino and requirements of young men.

Various investigators have pointed out that to determine the minimum amound of dictary protein necessary to maintain mitrogen equilibrium data are required for at least two levels of mitrogen intake near the region of balance in order that the value at exact equilibrium may be obtained by interpolation or extrapolation

Procedure ingest a nonnitrogenous diet as described above, insuring sufficient energy to provide for the body's requirements so that oxidation of tissue for this purpose may be avoided Feces and urine are collected as earlied in the early part of this chapter By means of kjeldahl analyses values are determined for metabolic nitrogen of feces and endogenous nitrogen of the urine Foliow this period with a test period during which the protein under investigation is included in the experimental diet. The foliowing

equation represents the percentage of the absorbed food ultrogen which is retained by the body

|Food N ~ (Fecal N ~ Metabolic N) ~ (Urinary N ~ Endogenous N)]i00 | Food N ~ (Fecal N ~ Metabolic N)

- Biological Value

3. Protein-sparing Action of Corbohydrate and Fat. The nonnitrogenous nutrients, earbohydrate and fat, have the power to diminish the extent of the catabolism of protein in the normal human body. In other words, they are said to "spare" protein. This point is illustrated in data reported by Yon Noorden and Dieters, which are tabulated below.

PROTEIN SI	PARINO	Action	OF	CARBOHYORATE
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Nutrogen Ingested	Netrogen en Urene	Remarks
12 6 g	10 4 g	
12 6 g + 200 g sucrose	9 0 g	i3 per cent reduction in protein entabolism

It will be observed that the addition of 200 g of sucrose to the diet was accompanied by a decrease of 13 per cent in the amount of protein criabolized. It has been established that carbohy drates are more efficient protein sparers than are the fats. For example, Voit found carbohydrate to produce a 9 per cent decrease in protein catabolism whereas fats produced only a 7 per ceot decrease.

Experiment. Ingest a uniform diet of known or estimated nitrogen content for a period of four days Collect and preserve the urine accurately (see p. 1044) in 24-hour samples and analyze the excretion of the third and fourth doys for total nitrogen On the fifth day add 200 g of sucrose to the diet. Analyze this utine also for total nitrogen Calculate your results and tabulate the data as shown in the table above

Did the sucrose influence the catabolism of protein in your body?

4. Influence of o High Caloric Nonnitrogenous Diet When an individual fasts, a certain amount of protein tissue is consumed each day of the fast. The destruction of such tissue is rather low on the first day because the glycogen stores of the body are being utilized to furnish the necessary energy. If an individual, instead of fasting, ingests a diet of high calorific value and very low in introgen the output of nitrogen in the urne of the third or fourth day will be less than on the third or fourth day in fasting. This is due to the fact that the body derives sufficient energy from the high caloric diet and there is less destruction of tissue protein than occurs in fasting. For a discussion of energy value of foods ew "Determination of Fiel Value of Food" below, and the table "Composition of Foods," in the Appendix

Experiment Ingest a high-calorie diet which is very low in nitrogen or actually nonnitrogenous A satisfactory diet may include sugar, butter, starch, cream, agar-agar, and water (For energy values see below and table, Appendix) Ingest such a diet for three days Collect the urlne in 24-hour perlods, preserve, and analyze it for total nitrogen, acidity, and ammonia. Note the low nitrogen excretion on the third day as compared with the nitrogen output of the third day of fasting. If so desired, you may (at some later date) fast for three days and repeat the above analyses for comparison.

Determination of Fuel Value of Food. When organic substances are oxidized or burned in the human body, they liberate a certain amount of heat. This calorific

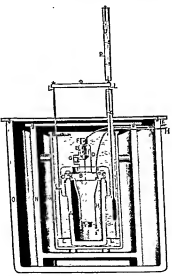


Fig. 260 Berthelot-Atwater Bomb Calorimeter.

energy or heat value varies according to the type of organic matter undergoing oxidation. Thus the proteins, fats, and carbohydrates of the diet when they are burned in the body yield different quantities of heat per unit of substance than do organic acids, alcohol, etc. The energy values of food protein, fat, and carbohydrate are as follows:

> Protein = 4 1 large Calories per g. Fat = 9.3 large Calories per g. Carbohydrate = 4 1 large Calories per g.

To allow for the meomplete digestion of these nutrients these coloric conversion factors are multiplied by the coefficients of digestibility and thus rounded off to 1, 9 and 4, respectively. The colorific value of foods may be estimated by applying these figures to the percentage composition is determined by chemical analysis. For these figures to the percentage composition is determined by chemical analysis. For ende fiber, and carbohydrate, the litter usually estimated by difference Crude fiber represents the insoluble, indigestible residue remaining after alternate extractions represents the insoluble, indigestible residue remaining after alternate extractions with and alkali, and consists chiefly of cellulose, etc., from cell walls. Such calculations are predicated on the assumption of complete availability of the carbohydrates lations are predicated on the assumption of complete availability of the factor 6.25 for consistentiated by difference, on the uon creal applicability of the factor 6.25 for consistentiate to a consist principally of glycerides.

In arriving at the total energy value of any given diet it is possible to burn weighed samples of the various foods in an only on a two-sphere to an apparatus called a bomb scalorimeter, illustrated in Fig. 200. By this means we may determine how much beat is calorimeter, illustrated in Fig. 200. By this means we may determine how much beat is calorimeter, illustrated in Fig. 200. By this means we may determine how much be made for the incompletely outlized substances of the urine and feees, e.g., organic nitrogen compounds. Thus while proteins yield about 5 T Calories per gram when burned in a compounds. Thus while proteins yield about 5 T Calories per gram when burned in a compounds reduces calorimeter, correction for incompletely oxidized urea and other N compounds reduces calorimeter, correction for incompletely oxidized urea and other N compounds reduces this value to the body to 4.1. Further correction must be made for indigestable carbon that the properties of the polysaccharides which in prior resist enzymatic digestion, is an accurate measure of the polysaccharides which in prior resist enzymatic digestion.

A simpler and less expossive form of apparatus for determining this calorific value of foods is the oxycalorimeter of Beoediet and Fox, in which the volume of oxygen required to burn a known weight of substacce is determined. A large mass of data required to burn a known weight of substacce is determined to burn a known weight of substacce is determined. A large mass of data required to burn a known weight of substacce is determined and tabulated, and it is therefore concerning the best value of foods has been collected and tabulated, and it is therefore possible to arrive at an approximate idea of the energy value of a diet by calculation (see table, Appendix)

5 Influence of Purme-free and High Purme Diets The urne and of the body has a twofold orgm, 10, 1t may arise from the metabols of the purme (nucleopro tem, nucleotide) material of body tissue (glindular organs in particular) or it may tem, nucleotide) material of body tissue (glindular organs in particular) or it may tem, nucleotide) material That urne and which arises from the first source is called endogenous while that which arises from the second source is termed source is called endogenous while that which arises from the second source is termed is still value Secretory activity may also act to merease the endogenous urne and output. The urne will therefore contain urne and even though no precursor of the and output. We may also increase the urne and output markedly by ingesting a high sungested. We may also increase the urne and output markedly by ingesting a light purne diet. However, no matter how much purne material is caten, only a small part purne diet. However, no matter how much purne material is caten, only a small part of it reappears in the urne as urne and That is, there must be some significant meta of it reappears in the urne as urne and That is, there must be some significant meta of the reappears in the urne and into other than exerction as urne and In gout there is an oble pathway for purne introgen other than exerction of urne and may be accumulation of urne and in the blood. In this disease the exerction of urne and may be accumulation of urne and increase considerably during an attack. The exerction of low before an attack and increase considerably during an attack. The exerction of exogenous urne and in gout is also much slower than normal

Experiment Ingest a purine free diet containing about 16 g of nitrogen and consisting of egg, cheese, milk, starch, fruit, sugar, and water for a period of two days (for purine content of foods, see table, p 1053) Determine or estimate the nitrogen content (see Appendix) and during the next two days substitute sweetbreads, thymus, or liver for all the nitrogen of the diet, maintaining the calorific value of the diet the same as before Return to the original purine-free diet for a third interval of two days During the final

period of two days feed a diet of sweetbreads or liver containing 50 per cent more nitrogen than that of the first sweetbread period Collect the urine for

INFLUENCE OF PURINE-FIRE AND HIGH PUBLIC DIETS NUTROGEY LYGESTION 10 G DAILY (TAYLOR AND ROSE)

Urinary Constituents Determined (g. per day)	Purine-free	Purine Diet	Purine Diet	Purine-
	Diel	(Medium)	(Increased)	free Diel
Uric acid N Total nitrogen Urea N (+NH ₄) Creatinine	0 09 8 9 7 3 I 57	0 14 8 7 7 1 1 49	0 24 9 1 7 1 1 51	0 07 8 8 7 05

each of the eight days of the experiment and determine pric acid and total nitrogen or urea. Note the rise in the uric acid output during the sweetbread periods. The uric acid output on the purine-free diet is endogenous in origin Tabulate your results. The data shown above were secured by Taylor and Rose in a similar but much more carefully controlled test than that just outlined

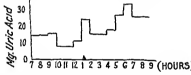


Fig 261 Influence of Protein Ingestion on Endogenous
Unic Acid Output Gluten (130 g) Ingested at 1 p m
C urtesy Mendel and Sighle J Bod Chem 22 215 (1915)



FIG 262 ENDOGENOUS URIC ACID OUTPUT DURING FASTING Courtesy Mendel and Stelle J Biol Chem 22 215 (1915)

6. A Study of Endogenous Uric Acid Output The uric acid in the urine originates from the purine material of the tissues (endogenous) and from the purine material specied (reogenous). Marcs claims that foodstuffs act to increase the endogenous ince acid output by stimulating the digestive glands to activity. A similar finding is experted by Mendel and Stehle The foodstuff having the most pronounced influence as protein Pilocarpina, which stimulates the digestive glands was found to increase the endogenous uric acid output whereas atropine which inhibits secretory activity.

was found to decrease the output of endogeoous une need. The influence of protein upon the endogenous une need exerction is shown by the chart to Fig. 261. The fasting output by the same individual is shown, for comparison, in Fig. 262.

PURINE CONTENT OF FOODS (AFTER BESSON AND SCHMID)

Food	Purine Aitrojen	Food	Purine Nitrogen
Meats Beef Ved Ved Mutton Pork Laver Tongue Sweetbreads Brains Fowl Chicken Goose Squab Fish Cod Salmon Herring Pike Trout Sardines Anchovies	per cent 0 037 0 038 0 026 0 011 0 093 0 055 0 330 0 028 0 029 0 033 0 058 0 021 0 060 0 047 0 056 0 118 0 145	Shellfish Oysters Crubs I obsters Vegetables Spinach Lentils Beans Mushroomy Peas Potytoes String beans Carrots Lettuce Cabbage Asparagus Cauliflower Fruits Bread Legg Cereals Butter Hull Cheese (except cream and dairy) Cream cheese Dairy cheese	per cent 0 029 0 020 0 022 0 022 0 025 0 017 0 018 0 002 0 002 0 003 0 003 0 008 0 00 0 00 0 00 0 00 0

Experiment Ingest a purine free diet consisting of milk, egg, fruit, cheese, butter, sugar, and bread for one day Continue the diet for breakfast and luncheon the next day but eat nothing after 12 o'clock noon, until 12 o'clock noon the following day, 1 e, the third day of the experiment At that time nigest 125-150 g of gluten or some other purine free protein preparation on the fourth day of the experiment eat nothing until 9 P M

Collect the urine each day in hour periods from 7 A M to 9 P M and analyze for uric acid (see methods on p 905, ff) Chart your data similarly to those shown in Figs 261 and 262, and compare them with the findings there recorded

The Rote of Purine Excretion The purine material ingested by the average normal person and not transformed in the body is eliminated in about 24 hours. In

the case of persons afflicted with gout the purine elimination is delayed. The establishment of this delayed purine elimination is often of diagnostic assistance.

Demonstrate the rate of purine excretion as follows. Ingest a purine free diet consisting of egg, milk, cheese, starch, sugar, fruit, and water for two days and follow this by a day in which sweethreads, thymus, or liver is substituted for one of the meals of the day (see table, p. 1053 for purine content of foods). Finish the experiment by Ingesting the original purine-free diet for two days. Collect each day's urine and analyze for uric acid flow soon after the sweethread ingestion was the nriginal plane of endogenous uric acid elimination reestablished? If one desires to locate this time more definitely the urine may be collected in short periods (I to 2 hours) and the uric acid content of each specimen determined. Particularly instructive data may be collected by berforming the above experiment on a gout patient and upon a normal person for comparison.

8. A Study of Creatinine Elimination. It has been established that a normal person ingesting a creatinine-free duct will exerte a uniform quantity of creatinine from day to day. The daily exertent of an adult man of average weight ranges from 1-1 og. For data as to creatinine exerction of a 60-kg man see Taylor and Rose s figures in the tablo op. 1052 The creatinine exerction depends primarily on the active mass of protoplasmic tissue, and therefore, it is generally frue that fat men will show a lower creatinine output than lean men of like body weight. The fact that, in a given individual, the creatinine output is uniform from day to day is made use of in metabolism experiments, for checking the completeness of 24 hour collections of unine. For further discussion of creatinine see p. 70.

Experiment Ingest an ordinary mixed diet (nonmeat) for a period of three days, varying the character of the duet dayly Collect the urine and analyze for creatinine (See p. 899 for methods of analysis)

Did the creatinine elimination change with the change in diet?

- 9 Formation of Hippuric Acid in the Human Body. Hippure and is present in human unnean small amount, about 0.7 g being excreted per day. The urns of herbivorous animals contains much larger quantities. This acid is formed in the liver in man, by the conjugation of benzon acid and glycine this formation is used clinically as a test of liver function. For procedure of the test see Chapter 31.
- 10 The Partition of Urinary Nitrogen and Sulfur as Influenced by Diet 14 was first shown by Folia¹⁴ that the percentage of the total nitrogen and total sulfur of the urine which appeared in the form of any particular nitrogenous constituent or in any particular form of sulfur was regulated directly by the extent of the total nitrogen and sulfur climination. This point is well illustrated in the table shown on p 10.5 which contains data regarding the so-called partition or distribution of the unusary nitrogen and sulfur.

It will be observed from an examination of this table that a normal protein diet which gave 16 8 g of urnary interiors yielded 87 5 per cent of this nitrogen as ures 3 per cent as a minimum a 30 fer cent as creatinum and 11 pr cent as ure card whereas nonprotein diet (starch and cream containing about 1 g, of introgen) which gave only 36 g, of urnary introgen yielded only 61 per cent of this nitrogen as urea but gave a girally increased percentage output in the case of each of the other nitrogenous constituints minimized [c] 113 per cent as a minimized. 25 per cent as urea and The percentage output of neutral sulfur was also greatly increased.

[&]quot; Folin 1m J I Ayetol 12, 118 (1305)

It will further be observed that the octual doily output of certain of the constituents is uninfluenced by the amount of protein ingested. Among these are creatinine and neutral sulfur. On the other hand, the output of morganic sulfur and uren is more or less directly proportional to the protein ingestion. The observation of such facts as these led Folin to formulate his theory of protein metabolism, which held sway for many years, but which has recently been considerably modified (see discussion on p. 1021).

THE NITROGEN AND SULFUR PARTITIONS AS INFIGURACED BY DIETAS

	Nort	nal Protein	Diet	Starch-cream Diet		
Constituent of the Urine	Amount, groms	Nitrogen, grams	Per cent of total V or S	Amount, grans	λ urogen, grams	Per cent of total N or S
Urea	31 6	14 7	87 5	4 72	2 2	61 7
Ammonia	0.6	0.49	30	0 51	0 42	11 3
Creatinine	1 55	0 58	36	1 61	0 60	17 2
Une acid	0 54	0 18	11	0 27	0 00	25
Undetermined N	1	0 85	49		0 27	7 5
Total N	1	16 8	100 0	Ì	3 0	100 0
Inorganic SOa	3 27	1	90 0	0 46		60 5
Ethereal SO:	0 19	1	5 2	0 10		13 2
Neutral SOa	0 18	1	48	0 20		26 3
Total SO:	3 04	1	100 0	0.76		100 0

Experiment During a period of two or three days ingest an ordinary mixed diet containing 100 125 g of protein (16-20 g of mitrogen) per day Collect the urine accurately in 24-hour periods (p 1044), preserve it, and analyze the urine of the second and third days for total nitrogen, urea, creatinine, total suifur, morganic sulfates, ethereal sulfates, and neutral sulfur (by difference) For methods of analysis see Chapter 31 Follow this period by a three-day period in which a diet of starch and cream having a similar caforffic value is ingested Analyze the urine for the second and third days as indicated above Calculate your results and tabulate as shown in the table above How did the change in the diet after the metabolism of nitrogen and sulfur?

In calculating the calorific value of a diet make use of the following values protein or carbohydrate, 4 Calories per g, fat, 9 Calories per g

11 "Metobolic Product" Nitrogen in Faces A certain quota of the fecal nitrogen is due to the presence of residues of digestive secretions epithelial cells, bac terns, etc. The nitrogen in these forms has been called metabolic product nitrogen.

To determine this form of nutrogen one method¹⁴ of procedure is as follows Ingest a nonnitrogenous diet for a period of two days. The diet may include desired quantities of storch, ercon, sugor, butter, uater, ond sadum chloride. About 15g. ogor-agor should be added to the diet to prevent constipation and to insure exacuation of approximately the normal quantity of feces (For the influence of agar agar, see Exp. 5, p. 1046.) To separate the feces properly, ingest a capsule of carmine at the beginning of the test and one of

⁴⁴ For a discussion of other methods of estimating metabolic product nitrogen see Forber Mangels and Morgan J Agr Research 9 405 (1917) and Schneider J Biol Chem 109 249 (1935)

charcoal at the end (see p. 1046) Preserve the feces as described on p 1046 After mixing the feces thoroughly, determine the nitrogen in weighed quantities by the kieldahl method, " according to directions given on p. 874 Calculate the quantity of netrogen eliminated per day. Inasmuch as no nitrogen was ingested, the natrogen present in the feces is of metabolic origin, i e , it is made up principally of nitragen in the farm of cells, digestive secretions, and bacteria

12 Influence of Defective Mastication on Food Residues in Feces. Rapid eat ing accompanied by defective mastication leads to the appearance of relatively large macroscopic food residues in the feces Uoder some conditions, however, protein utilization (see p 1047) may be as satisfactory after bolting of food as when it is very thoroughly masticated. This problem may be studied by the following method

o. Ingest a diet containing meat and be certain to masticate the diet very thoroughly. Collect a stool, examine macroscopically, mix carefully and

examine microscopically (see pp 448 and 449).

b. Ingest a diet similar to that employed in Exp o, above. Bolt the food, le, ingest it practically without mastication. Examine the feces as above Note the difference in the macroscopical and microscopical findings under (a) and (b).

if the nitrogen of food and feces to determined, we may calculate the protein utilization (see Exp 2, p 1047) By the additional determination of urlnary nitrogen, we may prepare a nitrogen balance (see Exp 13)

13. Preparation of a Metabolic Balance This test entails the analysis of the food ingested and of the urine and feces excreted, i.e. a study of the income and oulgo Proceed as follows

Select a diet which is simple, i.e., consists of few constituents, and which lends itself readily to accurate chemical analysis. A good type of diet for

BALANCE OF CALCIEM, MAGNESIUM PHOSPHORUS SULFUR, AND NITROGEN IN ACROMEGALY

	Calcium Ozide	Magnessum Oxide	Phosphoric Anhydride	Sulfur	Natrogen
			Grams		
Ingestion (daily) Lixerction (urine) Lixerction (feees) Exerction (total) Retention (daily) Rutention (per cent)	1 194 0 159 1 093 1 252 0 242 16 2	0 486 0 160 0 226 0 386 0 100 20 6	3 192 1 701 1 002 2 703 0 489 15 3	1 190 1 006 0 135 1 141 0 049 4 1	18 84 17 60 1 10 18 70 0 14 0 7

ordinary metabolism experiments of this sort consists of crackers (graham or soda), milk, butter, water, and agar-agar (to prevent constipation). Meat, especially prepared in quantity sufficient for an entire experiment, may also be utilized Ingest uniform quantities of these dietary constituents each day

[&]quot;In the outlation process use 10 g of prinsium sulfate instead of the copper sulfate The remail der of the procedure is the same as for urine

for a period of three days. Make an accurate collection of the urine passed during this interval (seo p. 1044). Separate the feces representing the three-day period (see p. 1046), and analyze foods, urine, and feces. The balances ordinarily prepared are those for nitrogen, sulfur, phosphorus, and calcium. Analytical methods for the determination of these elements may be found in Chapter 31.

The table on p. 1056 includes balances obtained in a metabolism test in a case of acromegaly.

14. The Influence of Woter on Metabolism. It has been demonstrated that increased water ingestion influences favorably many of the functions and activities of the human body is The increase in protein catabolism which accompanies high water intake is shown in the following data collected from an experiment upon a normal man. In this experiment the water ingestion at meals was increased 3 literaper day during the water period.

INFIUENCE OF HIGH WATER INTAKE UPON URINE VOLUME AND NITROGEN PARTITION

Day of Period	Urine Volume, ml	Nitrogen,	Urea Nitrogen, g	Ammonia Nitrogen, g	Crealinine Nitrogen, g	Creatine Nitrogen g
		Pro	eliminary Pe	rnod		
4	830	12 987	11 338	0 288	0 629	
5	920	12 084	11 476	0 305	0 619	ĺ
ß	880	13 183	11 568	0 369	0 651	
			Water Perio	d		
1	3440	14 161	12 596	0 486	0 610	0 063
2	3840	13 491	11 583	0 499	0 616	0 024
3	3670	12 981	11 212	0 553	0 589	0 102
4	3610	12 976	11 455	0 485	0 608	0 055
5	4020	13 138	11 879	0 456	0 589	0 128

The above data indicate an increased catabolism of protein material as shown by an increased output of total introgen upon the first and second days of the water period Part of this increase may, however, have been due to a flushing of the tissues rather than to increased catabolism of protein structures

[&]quot;Hawk. The relationship of water to certain life processes and more especially to nutrition" Read before discrete American Philadelphia (See Brocken, Bull., 3, 420 (1914) also Water as a Dretary Constituent' in Encorancing and Metabotican New York and London D Appletion and Co. 1924, Vol. 3, 327 (1923)) and all discrete New York and London D Appletion and Co. 1924, Vol. 3, 2157 (1923)) and latter by Underbill and Sallick, U Bull. Cham, 6,8,16 (1925) cannot be advanced and latter by Underbill and Sallick, U Bull. Cham, 6,8, 16 (1925) cannot be advanced argument against the value of a high water intake for the average individual. The observant mentioned experimented on animals and introduced as much as 50 mll water per kg body weight avery half hour until a definite toxic result was obtained. A similar excessive ingestion of any standard food would probably produce results fully as acrouss.

a RELATION OF WATER INTAKE TO VOLUME AND SPECIFIC GRAVITY OF THE URINE Ingest an ordinary mused diet for two days Collect the urine in 24 hour periods During the first day ingest very little fluid of any kind either at meals or between meals. On the second day ingest as much water as you can without physical inconvenience. A person of average size should have no difficulty in drinking 5-6 liters per day.

Measure the volume of each day a urine and take the specific gravity. Note the pronounced increase in volume and the low specific gravity of the urine

under the influence of high water ingestion

- b INFLUENCE ON PROTEIN CATABOLISM. That water stimulates protein catabolism may easily he demonstrated as follows: Ingest a uniform diet (milk, crackers, butter, peanute butter, and water) for a period of four days. During the first two days ingest your customary volume of water per day. During the last two days increase the water ingestion to 3-6 liters per day. Collect urine in 24 hour periods and analyze for total introgen by fieldabl method (see p. 874). Note the increased excretion of nitrogen under the influence of high water intake. If time permits, other nitrogenous urinary constituents may be determined (see table above).
- c INFLIENCE OF WATER DEFICIENCY. The importance of water in nutrition may be shown very satisfactorily in guinea pigs. Proceed as follows. Place two young pigs (150 20g.) In separate cages, and give each free access to a dlet of hay, oats and lemon or orange juice which has been dired rapidly at a low temperature. Permit one pig water ad lih, and give the second pig no water. The pig receiving water will remain normal and will eighbit normal gain in hody weight. The pig receiving no water will soon show pronounced losses in hody weight and other signs of abnormality. The animal will die ha short time unless water is added to the diet. This experiment demonstrates very clearly that water is an indispensable dietary constituent in fact, water is more important than food. The following experiment will show this.
- d Food Starvation vs. Water Starvation Place two young guinea pigs (150 200 g.) In separate cages Give one a dlet such as that described above plus orange juice (5 m l) but no water. Give the second pig no food, but per mit free access to water. The pig receiving no water will quickly become about main and it will be necessary to give it water to preserve its his. The second pig, which has access to water. But receives no food to eat, will live longer than the pig receiving an abundance of dry food. This little experiment impresses the important fact that man can live longer without lood than with out water. By restricting the amount of water in the diet of an albino rat the animal may be kept at constant body weight for several weeks although the diet is otherwise adequate. In many respects the effect as similar to that of underleeding. Some increased tolerance to water restriction may develop during such a test.
 - 15 The Metabolism of Fasting * The metabolism of a lasting man is entirely different firm their etabolism of a well nourshed person. The collection and analysis of it curned I ring a short fast (three to seven days) will demonstrate many important facts. The foll wing table with contains a data from factor tests made in the senior.

[&]quot;In case thepg Leanot donk the water the animal should be fed the fluid by a sound with the thermal in I fast it nee Benedet. A Study of Prolonged Pasting Carnege Inst. Wash. Pah. "33 1114 M. gibs Fast in your leadens with New York. I. Dutton and Co. 1023 and Jackson Inamison and Mainstrason Philadelphia, P. Blakiston a von and Co. Inc. 124.

author's laboratory, *1 illustrates some of the points in which fasting metabolism differs from normal metabolism

METABOLISM IN FASTING

Day of Period	Body H eight, kg	Total	1mmonta V, g	Creatine	Aculity ml O 1 N \aOH	P-O ₅	Chloride as NaCl,
		Pr	eliminary l	eeding Pe	rod		
1-4	Av 74 16	10 430	0 112	∖one	238 6	2 768	9 007
			I astını	g Period			
1 2 3 4 5 6 7	73 32 71 98 70 92 70 24 69 61 69 12 68 70	10 072 15 072 14 463 13 050 11 801 11 214 10 734	0 288 0 642 0 862 1 201 1 266 1 373 1 371	0 269 0 073 0 089 0 068 0 033 0 022 0 003	328 9 677 1 770 4 664 2 525 0 462 4 438 0	2 616 2 509 2 851 2 490 2 376 1 186 0 955	5 035 3 231 2 539 1 253 1 474 1 132 1 137

Abstinence from food for a few days can in no way operate to the disadvantage of a normal person. In fact, individuals affected with certain types of gastrointestinal dis-

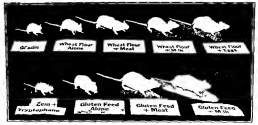


Fig 263 Showing Importance of Adequate Protein (Amino Acids) in the Diet

Courtesy Mendel Nutriton—The Chemistry of Life New Haven Yale University Press 1923 orders are benefited by fasting The fasting treatment has also been used in cases of diabetes mellitus and in the treatment of obesity

In order to determine experimentally how the fasting metabolism differs from normal metabolism, proceed as follows Ingest an ordinary mixed diet and collect your urine (see p 1044) for a day. Measure the volume and analyze the sample for total nitrogen, ammonia, creatine, sodium chloride, total

⁹¹ The chloride phosphate and acidity determinations were collected during one seven-day fast and the other data collected during a different fast on the same man

phosphates, and acidity¹¹ (for methods see Chapter 31) For the next few days (three to seven as desired) ingest nothing but water and collect the urios accurately and analyze for the constituents enumerated above Tabulate your results and compare them with those given in the table above

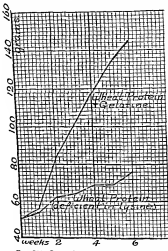


FIG 264 CURVE SHOWING INFLUENCE OF A DE-FIGURACT OF LIBINE IN THE DIET Lapublished data from the secure author a laboratory

16 Influence of Protein (Amino Acid) Deficiency. Certain of the amino acids which occur in proteins cannot be synthesized in the animal body. This subject is discussed on p. 101a, The importance of proper protein (amino acids) in the det is illustrated in F.g. 263. The following experiment which may readily be made using white rats as suljects will clearly demonstrate the importance of the amino acid lysine.

Demonstration of Lysine Deficiency. Place two young white rats (40-60 g.) in separate cages Feed one rat Diet 1 and the α the following table

^{*1} A more accurate experiment may be carried out by ngesting a uniform diet of known composition (see p. 104) for a few days before the fast

LYSIN DIRICILAGI DIET

		Diet 2 per cent
Rolled oats*2	60	60
Gelatin ⁹³	0	10
Dextrin or starch	30	20
Salt mixture	4	-1
Hydrogenated vegetable oil	5	5
Cod liver oil	1	1

The rat receiving Diet 2 will grow normally because of the high lysme content of gelatin The animal receiving Diet 1 will fall to grow properly because of lysine deficiency See Fig. 264.

III. MICROBIOLOGICAL DETERMINATION OF AMINO ACIDS

The widespread use of microorganisms for the assay of the vitamius in the B-complex group soon revealed their requirements for the annino acids Extensive investigations followed and successful quantitative assays for 14 amino acids, using bacteria, appeared shortly. At least three amino acids may be determined with mutant strains of the mold Neuroapora (see p. 1066), and one, 1-lysine, with a specific decarboxylase Of the ammo acids recognized as physiological constituents, only uorvalme, norleucine, and hydroxyproline have not been found essential for any organism yet investigated The fundamental principle involved in microbiological assays is to measure the response of bacteria, yeasts, or molds to graded increments of the sample and of a standard solution added to media furnishing all the nutrients required by the microorganism except the amino acid (or other nutrient) under assay The graded response may be measured by the increase in population of the microorganisms (i.e. turbidimetrically) or by their products of metabolism (acid or CO: production)

Microbiological methods are advantageous in that several amino acids may be determined in a single prepared hydrolyzate with the same uncroorganism with only slight modification of the basal medium. Only the natural or L-forms are biologically active, except in the case of aspartic acid, whose b- and L-isomers are equally available to Lactobacillus delivides. However, the L-forms are more expensive than the synthetic br-mixtures, so that the latter are used in the basal medium and in the standard series when available. When substituting one form for the other, or when the hydrochlorides are used, appropriate adjustments should be made in the amounts employed. Microbiological methods are exceedingly simple compared to the cliemical isolation procedures. Moreover, they are not subject to isolation losses since determinations are made directly on the hydrolyzates.

⁴² Oat protein is low in lysine. Gelatin is relatively high in this amino acid (see p. 122)

Careful attention must be given to proper methods of preparing hydrolyzates for assay Acid is known to distroy tryptophan and, in the presence of earbohydrate, tyrosine Enzymatic digestion or hydrolysis with barium hydroxide should be employed for tryptophan assays, since hydrolysis with sodium hydroxide has been reported to give erratic results Alkaline hydrolysis racemizes the amino acids, so that such treatment must be continued until the racemization is complete, and the values found multiplied by two Unless there is complete assurance that

AMINO ACID REQUIREMENTS OF VARIOUS LACTIC ACID BACTERIA*

Amino Acid†	Lacto- bacıllus arabınosus	Lacto- bacıllus cases	I acto- bacıllus delbrucksı	Strepto- coccus faecalis	Strepto- coccus lactis	Leu conosioo mesen- teroides
Arginine Histidine Isoleucine Leucine I ysine Methionine Phenylalanino Threonine Tryptophan Valine Aspartie acid Cystine Cilutamic acid Clycine Froline Serine Tyrosine	# - + + + + + + + + + + + +	++++++++++++++++++	+ + ± ± + ± + + + + +	+-++++++++++++	+ + + + + + + + + + + + + + + + + + + +	+++++++++++++++++++++++++++++++++++++++

^{*} Snell Conference on Amino Acid Analysis of Proteins New York Academy of Sciences

no amino acid loss occurs during the preliminary preparation of hydrolyzates, caution should be observed in applying such analytical data to the composition of the intact protein

The qualitative amino acid requirements of six lactic acid bacteria are shown in the table above These microorganisms have been employed in the quantitative determination of 14 amino acids. To facilitate the application of these methods a typical procedure is described below, and pertment data regarding the preferred methods of preparation of the hydrolyzates, the concentrations of the amino acids at half-maximum and at maximum growth, and the nature of the responses measured are

[†] The syn bol + indicates that the amino acid is essential ± that some growth but not 1945 maximum growth occurs in its absence and - that the amino acid is not essential

[‡] Asparagine and glutamine were essential for growth and supplied the organism with ti ese amino acida

MICHARIOTOCICAL DEFERMINATION OF THE AMING ACIDS

			Concent L 18	Concentrates is of L seemer				
Amino Acid	Agent	Mecrosquarenst	At Half Maximum Grouth	Maximum Growth	Measured		irsparation of Nedta and Dolaxis of Tests Verporation of Nedta and Dolaxis of Tests	tails of Tests
			Mucrogram	Maragrama per 10				
Arg nine	10 per cent 11Cl	Streplococus faecales		1 100	Aesdits	J Biol Chem		
Il studine	10 1 cr cent 17Cl	Lactobacellus cases	25	25	Turk idity	I Red Clam		
Boleueina	10 per cent HCl	Leuconotice mesenteroides P CO	200	35	Acidity	T But Clean		200
Lucine	10 per cent HCl	Streptococus facialis	.0:	325	Aesdits	3 3		154, 53 (1944)
Ly sine	10 per cent HCl	Streptococus facolts	281	25	Ac det	, Jo		152, 83 (1044)
Noth: n ne Let (Lilan ne	10 jer cont HCI	Streptococus farcalis Streptococus faecalis Lactobacilius delbrackis I DS	2225	5825	Acidity	J Blot Chem	166, 32 (1915)	
lire and Trait phant	10 ler cent HCl b N NaOlf or 6 N Ra/Om.	Leuconosioc mesenteroides P 60 Laciobacilus cases Sireplacacius faecalus Sireplacaccus faecalus	2554	88883	Andito	Prof	161, 613 (1645) 161, 613 (1645) 161, 511 (1943) 166, 35 (1645)	
Valine	(or enzyne di		4	2	Acidity			
Aspartic acid	10 per cent MCs	Lactobactive arabinous 17-5 Streplococcus faeralis Lactobactive delbrache 17-6	285	\$ <u>3</u>	Acidity	Brol		152, 83 (1944)
Serine	10 per cent HC	Lactobacillus arabinosus 17 6	Izg	2 2 2 3	Acidits Codits	J Biol Chem.	55	651 (1915) 273 (1945) 152 52 (1940)
Tyrosing	5 N NaOnt	Lactobaculus delbracks LDS Lactobaculus delbracks LDS	82	250	Aerdaty	J Biol Clem	155	,

The second control of the second from the American Type Collecton Weshington 8 D C under the following pumber, Streeters Receive No 1990 Prepared 1997 Accessful of the American metal-received from 1997 Replaced from the Streeters of the American metal-received 1997 Accessful to the second from 1997 Replaced from 199 Samples are hydrolyzed ten hours at 120° C

summarized in the table on p 1063 References are made to the original publications for further details 214

1 Microbiological Determination of Isoleucine, Leucine, and Voline (Method of Shankmon) " Principle Isoleucine, leucine, and valine are deter mined by measurement of the growth stimulation of Lactobacillus arabinosus Samples are prepared for assay by the microbydrolytic procedure of McMahan and Snell, 16 employing 10 per cent hydrochlone acid. Each amino acid is determined separately by omitting it from the basal medium

Preparation of the Somple Weigh into a pyrex test tube an aliquot of the sample containing 100 mg of protein Add 1 ml of 10 per cent hydrochloric acid the tube in an oxygen flame lleat for 10 hours in an autoclave at 15 lb pressure, or in an oven at 120° C Cool, open the tuhe, and wash out with approximately 95 ml of water Adjust with 4 N sodium hydroxide to pli 6 8 to 70 and dilute to a concentration of approximately 7 5 µg of L-valine, 15 pg of L-leucine, or 10 pg of L isoleucine per mi

Preparation of Bosal Medium The composition of the basal medium is given in the following table

COMPOSITION OF BASAL MEDIUM FOR THE ASSAY OF LEUCINE ISOLELCINE AND VALINE

(The amourts shown are for the preparation of 000 ml of basal medium)

Glucose	эg	L(-) Leucine	100 mg
Sodium acctate	3 g.	pt-Isoleucine	100 mg
Salt sol ition 1	5 ml.	pi-Valine	100 mg
Salt solution B	o ml	L(-) Cystine	00 mg
	əmg	pi-Viethionine	o0 mg
Adenine		L(-) Tryptophan	17 mg
Guanine	5 mg.	L(-) Tyrosine	17 mg
Uracil	o mg.		o0 mg
Thiamine	50 µg	DL-Phenylalanine	200 mg
Riboflavin	100 µg	L(+) Glutamie acid	100 mg
Sicotime act l	100 µg	DI-Threonine	100 mg
Biotin	0 2 µg	DL-Alanine	
Pyridoxine	50 μg	L(-) Asparagine or aspartic acid	
Pantothenic acid	ο 0 μg	L(+) Lysine	100 mg
p-Ammobenzoic acid	əμg	L(+)-Arginine	20 mg

For use in the assay tubes, omit from the hasal medium that amino acid r which analysis is being made Adjust to pli 6 6 to 6 8 with 4 N \aOli after

¹¹⁸ For discussion of additional microbiological assays o smino acids the reader is referred reviews by Sucil Ads in Protein Chem 2, 85 (1945) Schweigert and Snell Vatration bet and Rets 16 4J (1947) Dunn Food Technology 1 269 (1947) Barton Wright he Microbiological lesay of the Vitamin B Complex and Amino Acids New York Pitman ublishing Corp 1952

^{**} Shankman J Biol Chem 150 305 (1943)

[&]quot; Mc Mahan and Snell J Biol Chem 152, 83 (1944)

^{**} Reagent grade co centrated lydrad force acid solute a co tains approximately 35 per er t tydroef l rie acid

of Late at the appr x ate c neer trate na f the amino acids present from the comouts n of 1l e test material.

preparation. Stote in the refrigerator and arrange storage flask with a siphon so that the amount required for the day's work may be readily obtained.

The amino acids may be weighed out directly or may be prepared as stock solutions in water, containing the desired amount in a few ml. Where necessary, a little hydrochloric acid may be used to aid in solution. Store the stock solutions in the refrigerator.

Salt solution .1 contains 25 g. cach potassium monohydrogen phosphate and potassium dihydrogen phosphate, in 250 ml. water. Salt solution B has the following composition: 10 g. magnesium sulfate heptahydrate, 0.5 g. sodium chloride, 0.5 g. ferrous sulfate heptahydrate, and 0.5 g. manganese sulfate tetrahydrate, iu 250 ml. water. Add a few drops of hydrochloric acid to solution B to keep from precipitating.

Prepare stock solutions of adenine, guanine, and uracil, containing 1 mg. per ml. Solution is aided by heating in the presence of a few drops of hydrochloric acid. Store in the refrigerator, and renew at frequent

intervals.

Preparo stock solutions in water of thiamine, calcium pantothenate, and pyridoxine, containing 100 µg. per ml. Store in the refrigerator and renew at frequent intervals. Prepare the riboflavin stock solution in 0.02 N acetic acid, to contain 100 µg. per ml. Store as above, and in addition keep from exposure to light.

Stock solutions of nicotinic acid (100 µg, per ml.) and biotin (2 µg, per ml.) are prepared in 50 per cent ethyl alcohol. The stock solution of p-aminobenzoic acid (50 µg, per ml.) is prepared in glacial acetic acid, and stored in a dark-glass bottle.

Procedure. Carry stab cultures²⁰ of Lactobacillus arabinosus 17-5 on yeast extract-dextrose-agar (Difco) and subculture monthly. After transfer, incubate the cultures at 30° C. for 24 to 48 hours, and then hold in the refrigerator. Prepare the inoculum for the assay tubes by transfer from the stock culture to a sterife centrifuge tube of the complete basal medium, containing all the amino acids listed. Incubate the inoculum at 30° C. for 24 hours before use. Centrifuge aseptically and wash 3 times by suspending in sterife 0.9 per cent saline and centrifuging. Finally suspend the cells in 30 ml. of saline.

Assay. Pipet into pyrex test tubes 0, 0.25, 0.50, 0.75, 1.0, 1.5, 2.0, 3.0, 4.0, and 5.0 mi. aliquots of the test extract. Add to each tube 5 ml. of basal medum lacking the particular amino acid to be determined, and sufficient distilled water to make a total volume of 10 ml. Prepare a similar series of standards containing the amino acid to be assayed, employing a pure solution of the appropriate amino acid to place of the test extract. Optimal concentrations are 15 ml. Dutille or t.-leuciene per ml. or 20 4.0. Le-isoleucine per ml. Plug the tubes with cotton and sterillze in an autoclave at 15 pounds pressure for 15 minutes. After cooling to room temperature, add one drop (0.03 ml.) of inoculum to each tube and incubate at 30° C. for 72 hours.

Transfer the contents of each tube to a 125-ml. Erlenmeyer flash, using a constant volume of distilled water as a wash. Titrate the lactic acid produced with 0.05 N sodium hydroxide using hromothymol blue as the indicator.

^{*} American Type Culture Collection, No. 8014.

CALCULATION For each amino acid assayed, plot the titrations for the standard series in ml of 0.1 \ sodium hydroxide against \(\mu \) of the standard used From the appropriate curve estimate the amount of each amino acid assayed per tube. Calculate the concentration per ml of test extract. Determine the amino acid content of the test sample from the values obtained from not less than five of the tubes which do not vary by more than \(\pm \) 1 per cent from the average. Synthetic ni-mixtures contain 50 per cent of the biologically active enantomorph If if the racenie mixture was used as the standard, multiply the value obtained by one-half

In Fig. 250 is shown a standard curve obtained with μι-valine, during the assay of a β-lactoglobulin preparation for that amino acid. The method of calculation is illus-

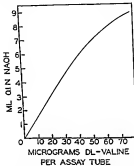


FIG 265 TITRATION OF I ACTIC ACID PRODUCED BY Lactobacillus arabinosus Grown at Different Levels of Valine Supplementation

trated in the table on p 1067. Results are expressed as per cent of value in the protein constituent of the material assayed.

There is some evidence that in addition to a proper balance of the essential amino acids it may be necessary to supply certain peptide linkages in the dictary to insure satisfactory utilization of protein intogen. Thus it has been demonstrated in rat assays that omes on of a new factor strepogenin from a dictary including a hydroly rate containing all of the essential amino acids in satisfactory quantities was respons the for a poor biological response. This factor believed to be a peptide is a growth stimulant for Lacidoscillus case and Streptococculacits. A preliminary method of assay has appeared based upon the growth of the former microorganism.

2 Use of Neurospora Mutants In 1941 Beadle and Tatum¹⁸⁰ reported the production of biochemical mutants in the red bread mold Neurospora by irradiation with ultraviolet and x-rays These mutants are unable to

Sprince and Woolley J Exp Med 80 213 (1944)
 Beadle and Tatum Proc Vol Acad Sci. 27 499 (1941) Tatum and Beadle 2nd. 28 234 (1942)

carry out certain specific chemical reactions which are normally possible to the unmutated or wild-type strain. For example, the untreated mold is able to synthesize all the components of protoplasm it needs (vitamins, amino acids, etc.) on a medium containing only sucrose, intrate, morganic safts, and botin. Various mutant strains have been obtained which are unable to grow without the addition of certain specific vitamins or amino acids to the medium, i.e., the strain has lost the ability it at one time

MICROBIOLOGICAL ASSAY FOR VALING IN A \$\textit{BLACTOGLOBULIY PREPARATION} (\sim 14.6 Per Cent)*

(0 107 g	was hydrolyzed	l and diluted	at 700	ml)
----------	----------------	---------------	--------	-----

Test Extract Added to Issay Tube	Titration after Incubation	Dic-Valine Equivalent Evaluated from Standard Curves	DL-Valine per ml of Test Extract
ml	nd 01 \ NaOll	μg	μg
0	0	0	
0 2a	07	51	20 4 (omit)
0.50	1 2	8 1	10 2
0 75	18	12 2	16 3
10	2 2	14 7	14 7
15	3 3	22 0	14 7 Average
2 0	4 5	30 3	15 2/ 15 3
3 0	6.5	40 5	15 5
4.0	7 9	62 0	15 5
50	8 6	72 7	14 5/

Calculation

 $153 \times \frac{1}{2} \times 700 = 5.74$ mg of L-value per 100 mg protein in sample

possessed to synthesize that particular vitamin or amino acid. It is believed that this loss of specific biochemical power is due to the loss of a single gene, which would ordinarily control the particular reaction which no longer occurs in the mutant

The implications of these remarkable findings are obviously widespread, not only in the field of genetics but also in many other ways, one
of which is in the field of microbiological assay. The Neurospora mutants
with specific nutritional defects can be used for microbiological assay of
amino acids and other nutrients just as has been described for the use of
certain bacteria. For example, Horowitz and Beidle¹⁰ have described a
coholineless struin of Neurospora, i.e., a mutant which cannot synthesize
choline and is therefore unable to grow without the presence of added
choline in the medium. These authors have described a method for the
microbiological assay of choline by the use of this strain. The procedure
is substantially identical in principle with that described on p. 1004 for the

 $^{^{\}bullet}$ The theoretical nitrogen content of β lactoglobulin is 15.6 per cent 0.107 g contains 100 mg of protein

²⁰¹ Horowitz and Beadle J Biol Chem 150 325 (1943) See Chapter 35 Choline

microbiological assay of amino acids, except that growth response is followed by drying and weighing the mold mycelium. Comparison is made in terms of a standard curve obtained with known amounts of choline. Methods have also been described for the assay of certain amino acids (methionine, leucine, arginine) by the use of the proper Neurospora mutants, and in general this versatile phase of microbiological assay appears to be just beginning in so far as future application is concerned.

Another contribution of this work is to facilitate knowledge concerning gene action, and the reactions controlled by genes. If, as has been suggested, each step in a biochemical reaction is under the control of a single gene, studies of the various mutants should permit a better understanding of the intermediate stages in biochemical reactions. For example, by this method of approach the progressive reaction ornithine \rightarrow circuline \rightarrow arginine, postulated by Krebs and Henselett in connection with urea synthesis in mammalian liver (see p. 1039) has been shown to occur in distinct gene-controlled steps in Neurospora. While this does not necessarily mean that the reaction occurs in this way, or even occurs at all, in higher cells, it at least indicates the possibility of it so doing. There is no doubt that further studies of this type will add significantly to our knowledge of the fundamental chemical processes of protoplasm.

IV. EXPERIMENTS ON CARBOHYDRATE METABOLISM

- 1. Hyperglycemia Produced by Carbohydrate Ingestion. The average glucose content of normal blood is somewhat less than 0 1 per cent. This is increased (byperglycemia) after the ingestion of carbohydrate food. The increase is noted more quickly after the ingestion of monosacthandes than after the ingestion of the more complex carbohydrates. After the ingestion of 100 g of glucose the increase in the sugar of the blood sometimes occurs in three imputes.
 - (a) INPLEYCS OF CIRCORE In the morning before breakfast, or at least three hours after breakfast, determine the normal augar content of your blood by means of some accurate micromethod (see Chapter 23). Ingest 100 g. of glucose dissolved in 250 ml. of water, and again determine the blood sugar level at intervals of 5, 15, and 30 minutes, and one, two, and three hours. (Plot a curve similar to the one shown in Fig. 266.) The urine may also be examined for sugar at intervals of one hour after the sugar ingestion.

Repeat the experiment on another day using 250 g, of glucose and compare the results with those obtained after the ingestion of 100 g. Explain your findings. If desired, this experiment may be combined with the one on "Alimentary Glucosuria," below.

(b) INFLENCE OF STANCE Repeat the experiment as given above for glucose except that 170 g. of white bread or 100 g. of starch made into a pastered are substituted for the 100 g. of glucose.

The experiment may be repeated as described above, using an increased amount of starch.

The various experiments may be conducted on patients suffering from diabetes must have a variable and instructive data collected. The alimentary hyperglycemia will generally be slower in reaching its maximum and will be more prolonged than in the case of normal subjects. In some instances after the diabetic has ingested

¹⁹³ In making starch pasts rub up the dry starch in a mortar with cold water, pour the suspended starch granules into boding water and star

 $100 \, \mathrm{g}$ of glucose the blood sugar does not reach its maximum until a period of two hours has elapsed. The blood sugar also returns to its former level more slowly than in the case of normal individuals.

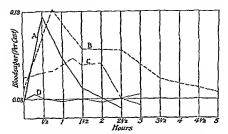


Fig 266 Blood Sugar As Influenced Bt Dier A, glucose, B, starch, C, starch and fat, D, fat

2. Experiments on Alimentary Glucosuria. According to Folin and Berglund, the sugar of normal urino consists of a motiey variety of carbohydrate products and carbohydrate derivatives including disand polysacchardes. These are believed to arise from foreign, unusable carbohydrate materials present in grains, vegetables, and firmts and from decomposition products due to cooking, canning, and baking of food. The ingestion of pure glucose, fructose, maltose, destrin, or starch does not normally give rise to glucinessis, but the ingestion of impure products or of an ordinary carbohydrate meal increases the sugar of the urine. So also may lactose and galactose.

Procedure. On arising at 7 00 AM, the student should empty his bladder and discard the urine voided. He should then drink one glass of water but eat no food. At 8 00 AM, the bladder is again emptled and the urine kept for analysis. The student then immediately drinks 200 g. of pure glucose dissolved in about 500 ml. of water. Urine specimens are collected again at 9 00, 10 00, and II 00 AM Test each specimen for sugar by Benedict's qualitative reduction test Determine sugar in each specimen according to the method of Folin and Svedberg (see Chapter 31).

Other students should go through the same procedure with the exception that for glucose there should be substituted 200 g. of came sugar, 200 g. of commercial destrin, 200 g. of pure destrin-starch, ¹⁰³ 100 g. of pure lactose, a dozen graham crackers with water only, two baked apples, bread and butter, meat with water only, or a large dish of pure gleatin without added signs.

This experiment may be made more complete by minking determinations of blood sugar at short intervals as described in Exp. i, p 1068. If desired, data on glucosuria, hyperglycemia, and carbohydrate in feces (see below) may be collected from one experiment.

Effect of Exercise on the Composition of the Urine. After strenuous exercise
the urine volume generally decreases, while the acidity and ammonia increase. The

^{**} Dextrue-starch mixture may be prepared by bothing 200 g. of starch in 900 ml of water cooling to 50° C, adding a hitle malt extract, and allowing to digest until completely inquefied

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phosphate and lactic acid excretion increase and the chloride exerction decreases The maximum effect usually is obtained in 20 minutes. Normal values should be restored in about an hour

Procedure. The subject who has had no food for several hours is given 50 ml of water every 15 minutes throughout the experiment, beginning for example at 12 00 noon At 12 00 the bladder is emptied and urine specimens collected every 15 minutes thereafter. At 1 00 the subject engages in strenu-

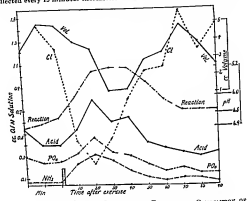


Fig 267 Curves Showing Influence of Exercise on Composition of THE URINE

Courtesy Wilson et al J Bud Chem 65, 753 (1924)

ous exercise for 2-3 minutes Running up and down stairs for this period will suffice In the urine specimens determine lactic acid, phosphate, total acidity, chloride, ammonia, and pli (For methods, see Chapter 31) Plot curves of amounts of these constituents per 10-minute specimen against time to show the effect of exercise on urine composition. The blood sugar may also be determined at intervals and the influence of the exercise noted Curves from a typical experiment are shown in Fig. 267

- 4 The Effect of Insulin on the Blood Sugar Level See the experiment described on p 769
- 5 Influence of Carbohydrate Deficiency Carbohydrates occupy a very promi eest place in the diet of man That they are not essential dietary constituents, at least for the white rat, may be shown by the following experiment

Demonstration on Carbohydrate Deficiency Use young white rats as subjects, feeding one rat Diet 1 and another rat Diet 2 as given in the following table

CARROHYDRATE DEFICIENCY DIET

	Diet 1 per cent	Diet 2 per cent
Casem Butter fat Lard Starch or dextru Yeast, dried, 104 grain per d 13	55 30 10 0 0 4	20 15 10 55 0 4

Both rats will grow normally in spite of the practical absence of carbohydrates in Diet 1 in the case of man, the withdrawal of carbohydrates is followed by ketosis (see p 1073) This ketosis is absent or much less pronounced in the case of the white rat

6 Isolation and Determination of Liver Glycogen. This exercise serves to illustrate the conditions for the manufacture and storage of liver glycogen by the animal organism, to provide some liver glycogen for study, and to show how tissue glycogen is quantitatively determined

Procedure Students may work in pairs Prior to the experiment, transfer a 6 mi portion of 30 per cent KOff to each of two 30 ml centrifuge tubes, stopper the tubes with a rubber stopper, and weigh on an analytical balance Suspend the tube on the balance by means of a copper wire fastened around the tube just below the pouring lip. Take care that the alkall does not come in contact with the rubber stopper.

Each group of four students will be supplied with a rat. Half the animals dispensed will have been starved and the other half fed. Remove a rat from its cage gently to avoid exciting it. Stun it by a blow on the head and de capitate it quickly Immediately remove the iner, weigh quickly to the nearest 0 1 g, and cut into two approximately equal portions, one for each pair of students. Mince the liver portion immediately and transfer samples of the minced liver to each of the two centrifuge tubes (0.7 to 1.0 g. samples from fed rats and 1 to 1.5 g. samples from starved rats). Stopper the tubes and weigh again. The difference between this weight and the original weight occurred the tube pius the KOH solution will give the weight of liver sample used. Remove the stoppers and place the tubes upright in a boiling water bath for 15 to 20 minutes, agitating the solution occasionally to ensure thorough disintegration.

At this point each pair of students should exchange one of the two tubes for one from another pair of students that represents a rat of opposite type. Thus each group will have one git cogen preparation from a fed rat and one from a fasted rat \(\text{dd}\)? mis \(\text{5}\) per cent \(\text{alcohol}\) to \(\text{each}\) tube, \(\text{mix}\) by \(\text{tapping}\), and immerse in the water bath until boiling just begins (care must be taken to avoid loss by sudden foaming). Allow the tubes to cool at room temperature for about 2 hours. Centrifuge, decant and discard the supernatant fluid, \(\text{drain}\) and wash the precipitates twice \(\text{with}\) 5-ml portions of 60 per cent alcohol by centrifuging, decanting, and draining as before \(\text{Expel}\) the

⁴⁴ The year is fed separately 04 g to each tat it may be fed in the form of a powder or as tablets. The only carbohy drate present in Diet 1 is the very small amount in the dred yeast.

last traces of alcohol by immersing the tubes in boiling water just long enough to dry the glycogen. Compare the relative amount of glycogen obtained from the fed animal with that from the fasted animal. To each tube add 10 ml distliled water and stir until a uniform suspension is obtained. Pipet 5 ml. of the suspension into a clean test tube, add 5 ml. 2 N sulfuric acid solution, insert a small funnel in the mouth of the test tube to minimize loss by evaporation, and heat in a boiling water bath for 3 to 4 hours to hydrolyze the glycogen Cool, add a drop of phenol red indicator, and neutralize cautiously with IN MaOII, with constant stirring, a volume of about 10 ml. should be required. Transfer the neutralized solution quantitatively to a 50-ml. or 100-ml. volumetric flask, dilute to volume with water, and mlx. Analyze this solution for glucose by the Somogyi-Shaffer-Hartman method (p. 571), using the volume of aliquot that will contain not less than 0.1 mg. or more than 20 mg. of glucose, calculated on the assumption that the glycogen content of the liver from the fed animal is approximately 10 per cent on a wet weight basis, and that from the starved animal is 0 2 per cent (it may be zero). Calculate the liver glycogen content as grams of glucose per 100 4 of liver, and report results to the Instructor. In your discussion of the experiment, briefly discuss the factors which influence glycogen storage and depletion. Use the remaining glycogen suspension which was not hydrolyzed for performance of the lodine test (p. 281) and the saliva test (p. 281).

V. EXPERIMENTS ON FAT METABOLISM

- 1. Fat Utilization This may be determined in a manner entirely snalogous to that used in the determination of protein utilization (see p. 1047) The fat may be determined by the baxon nethod (see p. 454). It is claimed when moderate amounts of fat see fed that the fat of the feces is largely independent of the diet. Therefore, in order to secure accurate information regarding the utilization of food fat, it is said to be necessary to determine the feed fat on a nonfat diet. There may possibly be a fat excretion from the intestine but much of the lipide of the feces is found in cellular structures (bacteria, cythichial cells, it is.) (See Chapter 21).
- 2. Fat in Feces. A normal adult will digest and absorb at least 90 per cent of the fat in the diet when the amount ingested does not exceed 100 g If the diet contains an excessive amount of fat, e.g., 300 g per day, considerable appears in the feces In pancreatic diseases and such conditions as are accompanied by a decrease in bile flow, the digestion and assimilation of fat is leasened.

Experiment. (a) Ingest an ordinary mixed diet containing an average amount of fat per day, e.g., 75-100 g. Collect a stool and examine it microscopically as directed on p. 449 (b) Now logest a diet containing an excessive quantity of fat, e.g., 300 g. per day. Separate the feces and subject a representative sample of the feces from the high fat diet to microscopical examination (c) If it is desired, the fat may be extracted from some of the stool by applying the priociple involved in the quantitative determination of fat lot the Saxon method (see p. 454). Evaporate the ether extract and identify the fat in the residue by tests given in Chapter 3.

3 Influence of Fat (Fatty Acid) Deficiency. Although it has often been demonstrated that fats are not essential constituents of the diet from the energy standpoint, Burr and Burr have shown the essential nature of certain unsaturated fatty acids five discussion on p 1004). The following experiment demonstrates the characteristics of a deficiency of essential fatty acids in the rat.

Experiment. Two groups of rats, weaned at three weeks, are placed in individual cages and fed the following diet: purified casein105 24, sucrose 72.1. salt mixture (McCollum-Davis, see Appendix) 3.9. Supplementing this dlet, 0.65 g, of dried ether-extracted yeast upon which is dried 2 drops of an ether extract containing the nonsaponifiable fraction of 70 mg, cod-liver oil. Is fed dally. To each rat in one group 10 drops of lard are given daily. Water is supplied ad lib., and if desired, the quantity consumed may be measured.

In about 70 to 90 days the symptoms of the deficiency will begin to appear in the fat-free group. The legs, especially the hind legs, become scaly and swollen, the tail becomes spotted and ridged and finally necrotic (Fig. 268), the hair on the back becomes filled with dandruff and tends to fall out, and degeneration of the kidneys sets in, as may be observed from the bloody urine. The renal lesions are the immediate cause of death. In the females, ovulation becomes irregular and finally ceases.



FIG 268. NECROSIS OF THE TAIL OF A RAT ON A FAT-FREE DIET. Courtesy. Burr and Burr. J. Biol. Chem , 82, 345 (1929).

The deficient animals eat the same amount of food but drink twice as much water as the controls. Before the deficient animals reach a moribund condition, their diet may be supplemented with a small amount of an oil or fatty food, to test the curative properties of the fatty acid molecules.

The above procedure can be employed quantitatively for the bioassay of the essential fatty acids present in foods. 10% Weanling male rats from litters of which the mothers have been on a low-fat dlet are placed on a fat-free dlet until depleted, as determined by a constant body weight for 3 weeks. This requires a period of approximately 12 weeks. The rats are then divided into the following groups: negative control group; groups to determine reference growth curve (which receive 10, 20, or 50 mg. methyl linoleate daily); groups to receive unknown fat at several levels (25 and 50 mg. per day for cottonseed and similar oils, 250 and 500 mg. per day for margarines and butter). The assay period is continued for 8 weeks. The potency of the unknown fat can then be estimated from the log dose; gain-in-weight relationship, as is done in the standard procedure for bloassay of vitamin A.

4. Ketosis. Ketosis may be induced in a normal person by the ingestion of a carboby drate-free diet. The ongin of Letosis is discussed on p 1007. The intensity of Letosis

¹⁰⁵ Reprecipitated curd casein is washed until free from chlorides, extracted repeatedly. with alcohol, and finally extracted for one week with ether, and dried at 35° C For detailed directions see Burr and Burr J. Biol Chem., 82, 345 (1929).

134 Deuel, Greenberg, Anisfeld, and Melnick J. Nutrition, 45, 535 (1951)

may be evaluated by determination of the excretion of ketonic bothes in the urner (ketoniura). The following table shows the data obtained in an actual case of the withdrawal of earbohy-first food from the det of a normal man (to a Noorden).

KLTOSIS ACCOMPANYING CARBOHYDRATE WITHDRAWAL

Oay	Diet	Fxcretion of Acetone Bodies Cal- culated as β Hydroxybutyric Acid
		grams
1	Protein, fat, and carbohydrate	None
2	Protein and fat	0.8
3	Protein and fat	1 9
4	Protein and fat	8 7
5	Protein and fat	20 0
6	Protein, fat, and carbohydrate	2 2

Experiment Ingest an ordinary mixed diet for I day Follow this by a period of 2 to 4 days in which no digestable carbohydrate is eaten (A diet of meat, eags, butter, agar-agar, and water has a very low digestable carbohydrate value) Collect the urine for each day of the experiment, examine it qualitatively for actone bodies (see tests, Chapter 29) If present, determine the total acetooe bodies quantitatively (for methods see Chapter 31) The blood may also be examined (see Chapter 23) Did the withdrawal of carbohydrate food cause a ketonutia?

VI. ADEQUATE vs. OPTIMAL NUTRITION; THE DIETARY EFFICIENCY OF MILK

The foregoing experiments have demonstrated that certain factors must be present in a diet if it is to be deemed adequate But such a diet although it qualifies under all required nutritional standards, may be improved upon We thus form what may be termed an optimal diet. To demonstrate the difference between an adequate and an optimal duet proceed as follows

Place two white rats from the same litter, each weighing 35-45 g, in separate cages Feed one rat an adequate diet (see below) and the other an optimal diet (see below)

Adequate Diet		Optimal Diet		
Whole wheat Whole milk (powder) Sodium chloride		Whole wheat Whole milk (powder) Sodium chloride	66 33 1	

it will be noted that dry milk constitutes only one inth of the adequate det whereas it makes up one-third of the optimal diet. Give the animals water as lih and weigh them at least twice a week. Continue the experiment for at least 10 weeks, plotting the growth curve of each animal and keeping an accurate record of the food eath (see Appendix for methods of recording data) If it is desired to investigate the relationship of these diets to reproduction two rats of opposite sex may be used in each test and caged together

The rat (or rats) receiving the optimal diet should grow more rapidly than the animal or animals receiving the adequate diet. If the experiment embraces the question of reproduction it will be found that the rats on the optimal diet have greater success in

the rearing of young and that the offspring grow better during the missing period than do the offspring of the rats ingesting the adequate diet

In a comprehensive series of observations on these diets by Sherman and associates it has been found that the animals fed the optimal diet mature earlier and live longer in common with other data secured from tests with albino rats these findings may be applied directly to the human dietary, and afford a very striking demonstration of the remarkable nutritive efficiency of milk

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14, 1 (1945)

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Inorganic Metabolism

Composition of the Animal Body. The elements that compose the human organism and their approximate relative amounts in the body are as follows:

	Per Cent	Approximate Amount in a 70 kg Man		
		grams		
Oxygen	65 O	45,500		
Carbon	18 0	12,600		
Hydrogen	10 0	7,000		
Nitrogen	3 0	2,100		
Calcium	1 5	1 050		
Phosphorus	10	700		
Potassium	0 35	245		
Sulfur	0 25	175		
Sodium	0 15	105		
Chlorine	0 15	105		
Magnesium	0 05	35		
Iron	0 004	3		
Manganese	0 0003	0 2		
Copper	0 0002	0 1		
Iodine	0 00004	0 03		

Various other elements are also present in traces, these are sometimes called as a group the trace elements. In many instances their nutritional significance if any is obseure. In addition to copper, mangauese, zure, and iodine, which may be considered trace elements of known nutritional significance, this group includes the elements aluminum, fluorine, silicon, hithium, bromine, arsenie, lead, molybdenum, vanadium, and possibly others. Each of these elements is in fact a mixture of isolopes, elements that have identical chemical properties but differ in certain physical properties (see Chapter 32, "Isotopes")

The elements which make up the animal body are obtained from various sources. Carbon, hydrogen, and oxygen are supplied to the body by carbobydrates, fats, and proteins, and by water and the oxygen of the air, introgen, as well as a considerable proportion of phosphorus and sulfur, are supplied by proteins, the remaining elements are supplied by the mineral constituents of initial foodstuffs, by common salt, and to some extent by the minerals of druking water.

1077

1078 About 90 per cent of the total oxygen of the annual body and about 70 per cent of the total hydrogen are present together in the form of water, which makes up roughly two-thirds of the total body weight. The remaining oxygen and hydrogen, along with all the introgen, most of the carbon

PROXIMATE CHEMICAL COMPOSITION OF AN ABULT HUMAN BODY

Adapted from the data of Mitchell, Hamilton, Steggerda, and Bean $(J \; Biol \; Chem \; ,$ 158, 625 (1945)) The subject was an adult male, aged 35, estimated to have been in reasonably good nutritional state at the time of death, which was due to an acute heart attack The body was obtained from the Department of Anatomy of the Um versity of Illinois College of Medicine, and had been preserved only by freezing until the dissection for analysis was begun about 6 weeks after death Approved analytical methods were used

Parts Anal jzed	Per Cent of Total Body	lV ater	Lther Soluble Solids	Crude Protein (N × 6 25)	Ash	Calesum	Phos- phorus
Skin Skeleton Teeth Strated muscle Brain, spinal cord, and nerve trunks Liver Heart† Lungs† Spleen Kidneys Panereas Alimentary tract Adipose tissue Remaning tissues Liquid Solid Contents of alimentar tract Bile	0 8	50 09 93 3 70 4	3 35 12 68 10 35 9 26 1 54 1 19 4 01 3 13 08 7 0 2 9 42 4	15 88 13 38 17 81 14 60 3 12 69 4 13 10 4 7 00 7 5 68	0 95 1 13 0 90 0 03 0 80 0 51	0 0143 0 0125 0 0116	0 113 0 114 0 217 0 174 0 15 0 111 0 044
Contents of ahmenta	0 8 0 1 0 0	5 3				1.	_

^{*} Assumed † Somewhat enlarged

and sulfur, and some of the phosphorus are found in the organic compounds of the body (carhohydrates hpides proteins, etc.) These compounds comprise about 90 per cent of the total solid matter of the animal body, the remaining 10 per cent (about 33 per cent of the total body weight) being largely morganic The morganic elements essential for

^{\$} Somewhat cor gested

runn il life meliide sodium, potrissium, calcium, magnesium, phosphorus, iron, copper, chlorine, iodine, cobalt, manganese, zine, and possibly others

The relative amounts of water, organic material, and inorganic elements ("asb") chriateristic of the body as a whole do not necessarily represent the composition of the individual tissues and organs of the body, where wide variation may be encountered, not only from tissue to tissue but for a particular tissue under varying normal and pathological conditions. These differences between the various parts of the animal body to this respect are illustrated by the data in the table on p. 1078.

The term "proximate" refers to an analysis in terms of fundamental components but not necessarily in terms of the elements themselves Fresh tissues or tissues from different animal species may vary more or less from the values given in the table. The weight of the ash is slightly higher than the actual weight of mineral elements present, including as it must the carbon and oxygen of carbonates, the oxygen of phosphates and sulfates, etc.

The absence of carbohydrate from the table is to be noted. If the carbohydrate content is estimated by difference, it cannot account for more than a few per cent of the total body weight. This emphasizes the fact that io animal tissues as a whole, carbohydrate is present in limited amount, despite the high carbohydrate content of the animal diet—in contrast to place tissues, where carbohydrate may make up the major portion of the solid matter present Acother factor which enters into the picture is that much of the carbohydrate in animal tissues readily undergoes post-mortem changes, i.e., glycogen and glucose are converted to lactic acid, etc., so that a precise estimation of carbohydrate content requires special technical treatment.

WATER

Water is the most abundant compound in the animal body, making up approximately two thirds of the total body weight, although the various tissues within the body may differ significantly to water content (see table, p. 1078). Other organisms, e.g. the jelly-fish, may have a much higher proportion of water. Water furnishes what has been called "the aqueous milieu within which life processes occur." Water is more essential than food in the sense that an individual can survive much longer when deprived of food than when deprived of water According to Thorpe, death usually results when about 20 per cent of the body water is lost. For experiments illustrating the need for water in the diet, see p. 1052.

Water possesses a number of physical and chemical properties which promote its physiological utility. The high solvent power of water permits the formation of a variety of true and colloided solutions, within which reactions may occur at a much more rapid rate than if a solid phase were concerned. The high dielectric constant of water promotes iomizatioo, thus facilitating reaction occurrence and velocity. Water itself enters into a great many of the reactions of biological material (hydrolysis, ouddation, reduction, etc.) and appears to catalyze other reactions in which it plays no apparent part. The lubricating action of water is important in coo-

Chap 34

nection with such physiological functions as swallowing and the role of the various internal fluids of the body (synovial, peritoneal, pleural, etc.)

Water is a good heat insulator, and at the same time appears to be of major importance in the control of heat loss from the animal body. The heat required to convert I g of water to water vapor at room temperature is approximately 0.6 Cal. Du Bois found in one instance that the amount of water lost by vaporization from the skin and lungs of a noimal resting man under ordinary conditions of temperature and humidity was about 680 g per day. The heat loss associated with this vaporization represented about one-quarter of the total daily heat loss other means of heat loss include radiation and conduction, as the environmental temperature rises toward that of the animal body (37° C), these latter means become relatively ineffective and heat loss by vaporization of water becomes proportionately greater.

The significance of water in the animal body! has two general aspects (1) the relation between water entering the body and that leaving it, 10 the water balance, and (2) the distribution of water between the various water-containing fluids of the body

Water Balance. The sources of body water include (1) the fluids of the duct, (2) the solids of the duct which contain more or less water, and (3) the water produced by the oxidation of material within the ussues Tho major channels of water exerction include (1) tho lungs, (2) the skin, (3) the kidneys (urinc), (4) the intestinal canal (feces) in the lactating female the milk and (6) to a minor extent the tears. The relative significance of these various factors may be illustrated by the following table.

WATER BALANCE OF A NORMAL INDIVIDUAL

Source			}	Excretion		
	Ml	Per Cent of Total		мі	Per Cent of Total	
Fluids of diet	1200 1000	48 40	I ungs Skin	500 500	20 20	
Oxilation within	300	12	Urine Feces	1400 100	56 4	
Totals	2,00	100		2000	100	

Normally an individual is in water equilibrium 10 water gain equals water loss with respect to all sources A water intake in excess of the al-lity of the body to excrete water may be toxic (water intoxication), a water loss which exceeds water intake may lead to dehydration, and if sufficiently prolonged to death

For a review see Mbott Am J Med Sci. 211–232 (1946) See also Hawk. The relational profit water to certain He processes and in receive in 1940 to intuition. Biochem Biolium 3, 420 (1944) Hawk. Water as a Detary Constituent in Endocrinology and Mediclium VI. 111 Lo. Ion and New York. D. Applit in and Co. 1924.

It has been demonstrated that inorganic elements must be available in the animal diet not only in adequate amount but also in proper balance with respect to one another. For example, a high intake of potassium salts results in increased elimination of sodium and chlorine, adequate dietary calcium is necessary for retention of potassium, and in cases of vitamin D deficiency, too high or too low ratios of ealcium to phosphorus in the diet lead to serious impairment of bone structure.

In the following pages are considered some of the metabolic aspects of inorganic elements and compounds. Quantitative methods for the determination of various inorganic elements in blood and urine have been described in Chapters 23 and 31; the occurrence and significance of these

elements in the urine are discussed in Chapter 28.

SODIUM

This element is present to a considerable extent in the diet and in the body fluids of the animal organism, invariably in the form of the sodium ion, Na⁺, and in the diet at least, for the most part as sodium chloride. However, the sodium ion is a physiological entity whose function within cortain limits is relatively independent of whatever anions (chloride, bicarbonate, lactate, phosphate, proteinate, etc.) are associated with it.

The adult human body contains about 100 g. sodium, distributed almost entirely in the extracellular fluids (blood plasma, interstitial fluid) of the body. On the average, about 4 to 5 g. sodium are ingested per day in the ordinary diet, corresponding to about 10 g. sodium elloride, which is the major form in which this element is ingested. Practically all of this ingested sodium is excreted by way of the urine. On diets low in sodium the urinary excretion falls to a very low level, so that the daily sodium requirement is considerably in excess of the amounts ordinarily ingested, and is commonly covered by the amount of sodium chloride (table salt) in the diet.

Carnivorous animals receive adequate salt from the animal tissues which constitute their food. The food of herbivorous animals is relatively low in sodium, and is high in potassium, the ingestion of which causes an increased urinary excretion of sodium. This loss of sodium induces a craving for this element, which in nature may be satisfied by such animals traveling long distances to salt lieks, while domestic cattle are furnished with salt blocks. In the case of human subjects, the addition of salt to the diet makes it possible to eat with relish larger amounts of vegetable foods. Lactating cows not given a salt supplement yield milk of low sodium chloride content and may suffer otherwise. It has been shown in rat experiments that sodium deficiency unfavorably affects the appetite, the normal increase in weight, the storage of energy, and the synthesis of fat and protein. On a diet otherwise normal hut of restricted sodium content (0.002 per cent), rats show retarded growth and disturbances of the eyes and reproductive function, with death ultimately ensuing. Rats may have their sodium needs satisfied by diets containing as little as 0.1 per cent of sodium chloride.

It has been shown by the use of radioactive sodium that this element is absorbed exceedingly rapidly from the gastrointestinal tract, some absorption apparently taking place even from the stomach—somewhat surprising in view of the relatively restricted absorptive capacities of this organ. Radioactive sodium (Na2) has also been used as a means of determining the volume of extracellular fluid in the animal body, on the assumption (which appears to be reasonably valid) that the sodium ion is distributed evenly or in a predictable way throughout the extracellular fluids of the organism and is virtually absent from the body cells.

The major functions of the sodium ion in the animal body appear to be in connection with osmotic-pressure regulation and acid-base halance, although other possible functions such as a catalytic effect on enzyme activity, though as yet undiscovered, cannot be evaluded. The role of sodium in osmotic pressure regulation and acid-base balance is hest uoderstood by a consideration of the electrolyte distribution of a typical body fluid (blood plasma) as shown herewith.

APPROXIMATE ELECTROLITE DISTRIBUTION OF NORMAL HUMAN BLOOD PLASMA

HUMAN BLOOD PLASMA

"Bases"

"Acids"

Ion: Na + K + Ca + Mg = Cl + HCO₂ + Pr + Others

Concentration:
$$154 + 5 + 5 + 3 = 106 + 28 + 17 + 16$$

(milliequivalents
per liter of 167 = 167

plasma water)

The numerical values are to be considered as approximations only, subject to more or less wide variation under both normal and pathological conditions. As a matter of fact, the ionic concentration of an element io a biological fluid is somewhat difficult to define with exactness, since what is ordinarily measured is the total amount of element rather than that fraction which is ionized (see discussion of calcium, p. 1088). Nevertheless, the relationship shown above serves to bring out several fundamental facts.

In the first place, it must be true that the sum of all the cations present (Na, K, etc.) equals the sum of all the anions (Cl, HCO2, etc.). This is evident only if concentrations are expressed in terms of milliequivalents rather than milligrams. This is one reason for the increasing use of the former term in connection with such substances as sodium, chloride, bicarbonate, etc., in their relatinn to physiological processes. For example, if bicarbonate is replaced by chloride in plasma, this replacement is on the basis of ion for ion (i.e., equivalent for equivalent) rather than gram for gram. The equivalent concentration of a monovalent ion may be obtained by dividing the weight of inn present by the equivalent (combining) weight of the ion; far milliequivalents, both weights are expressed in milligrams. Thus if sodium is present in the extent of 354 mg. per 100 ml., or 3,540 mg. per liter, the milliequivalent concentration per liter is 3,540/23 = 154 milliequivalents (mEq). For a divalent ion such as Ca++, the equivalent weight is the combining weight divided by 2, since one Ca ion is equivalent to two monnyalent anions.

From the data on electrolyte distribution, therefore, it can be seen

that the sodium ion is the chief cation of blood plasma, and this is true also for the other extracellular fluids of the body. Within the cells, sodium is present in relatively low concentration, being replaced largely by potas-

sium and magnesium. The division of the various ions of plasma into "bases" and "acids" deserves some clarification, since this usage is rare in general chemistry though quite common in physiology and medicine. The plasma (and the tissue fluids in general) are essentially neutral in reaction, and remain so under all conditions compatible with life. When nn acid such as lactic acid enters into or is produced within a neutral buffered physiological fluid, to all intents and purposes the hydrogen ion of the neid disappears in the formation of water or other un-ionized molecule. The fluid still remains essentially neutral in reaction, and the only evidence that acid has been present is found in the increased amount of the acid anion-e.g. the lactate ion in this instance. This increased acid anion content is at the expense of such buffer anions as bicarbonate, phosphate, proteinate, etc.. originally present. Thus the entrance of acid is reflected not by any marked change in pH, but by alteration in the relative amount and distribution of the various anions present. The anions therefore represent the "acid" portion of the electrolytes An analogous situation prevails with respect to base, the entrance of which, within the framework of a neutral solution, will he evidenced almost entirely by an increase in cation (e.g. Na) content. Naturally, other factors enter into acid-hase balance within the body, as is evident from the discussion in Chapter 24, but the concept of acid-hase halance reflected in part by the kind and distribution of anions (acids) and cations (bases), within an essentially neutral solution, is of considerable value in an understanding of all phases of this subject. From what has been said, it can be seen that sodium is the chief base of the plasma, and of the extracellular fluids in general.

Another aspect of sodium-ion conceutration which is of importance is in conucction with osmotic pressure regulation. The total osmotic pressure of a hiological fluid such as the blood plasma is equal to the sum of the osmotic effectiveness of all the ions present. Thus the plasma corresponds osmotically to a solution 0 167 M (167 milliequivalents per liter of plasma water) with respect to an electrolyte such as NaCl, which is assumed to be completely dissociated in solution to yield two ions, or to a solution of a nonelectrolyte which is approximately 0.33 M (2 \times 0.167). The osmolar concentration of plasma is therefore approximately 0.33. Note that of the total osmotically effective bases of plasma, sodium makes up approximately 92 per cent (154,67). Thus changes in the osmotic pressure of plasma (and in general of the extracellular fluids of the body) are largely due to, and may be caused by, changes in the concentration of sodium present. It is true that under stress a loss of sodium may be compensated for to a small extent by an equivalent increase in potassium, but the ability of the organism to substitute bases in this manner is relatively restricted (in contrast to the wide variation in anion distribution which may occur) and any major loss of sodium from the body, as in Addison's disease, and prolonged diabetic ketosis, leads to a significant lowering of the osmotic pressure of the body fluids, and therefore water loss

(dehydration) Restoration is not complete until both the lost base and the lost water are replaced

The role of sodium in acid-hase belines is secondary to its role in maintaining the total osmotic pressure of the body fluids. That portion of the body sodium equivalent to the bicarbonate present (i.e., about 28 milliculus alents per liter of plasma water) represents most of the available base of the plasma which can be used for the neutralization of entering acids, and in conjunction with earbonic acid determines the pH of blood (see Chapter 24). It is more correct however, to regard acid-base changes in terms of the anions present rather than in terms of sodium. Thus one may have an acidosis or alkalosis in terms of variation in plasma bicarbonate and CO₂ tension without significant changes in sodium content.

The sodium ion has an effect on irritable tissues such as muscle which does not appear to be related to esmotic forces, and which is counteracted by the presence of the calcium ion, in the proportion of about 1 to 2 calcium ions per 100 sodium ions This is the hasis for the use of such "physiologically balanced" solutions as Ringer's, Tyrode's, etc., rather than isotonic saline alone, in experiments involving the maintenance of function of isolated animal tissues or organs

The excretion of sodium has been discussed on p 963 On an ordinary diet urnary sodium amounts to about 3 to 5 g per day, corresponding to 10 to 12 g expressed as sodium chloride, but wide variations naturally can be expected On a low salt diet and in starvation, urnary sodium chloride excretion falls to very low levels. The activity of the antidiuretic hormone of the posterior pituitary (see p. 775) is sensitive to the availability of corticosterione and deoxycorticosterione, elaborated by the adrenal corter. As a result, a deficiency of these steroid hormones is followed by a loss of sodium and chloride in the urne and retention of potassium, although the causative relationship has been questioned. The restriction of dietary sodium chloride is widely advocated in essential hypertension and cardiac disease.

CHLORINE

This element is found in biological material exclusively in the form of the chloride ion, "the biological significance of postulated organic chlorine compounds, such as chlorolipides, has as yet to be conclusively demonstrated. In the diet chloride is found largely as sodium chloride, the daily intake approximating 6 to 7 g, representing about 10 g of sodium chloride, but wide variation in this respect is encountered, depending as it does upon dictary habits. The adult human body contains approximately 100 g of chloride, found largely in the extracellular fluids of the body, but present to some extent in red blood cells and to a lesser extent in the other cells of the tissues.

Although a close relationship exists between chloride and sodium in

⁴ Fourman Perferences Keyler Denp-es Bartter and Albright Melabolism 1 259 (1952)

^{*}A notable exception to this rule is the antibiotic Chloromycetin(chloramphenicol) which contains chlorine in organic combination

certain physiological processes, the chloride ion has a number of functions which are peculiar to it and which are essentially independent of sodium ion functions. Thus it is the chief anion of the gastric juice, being present there in approximately the same concentration as in the blood, at least before dilution or other modification of the gastric juice, and accompanied by the hydrogen ion in substantially equivalent amount, rather than by sodium as in the plasma and extracellular fluids of the body. The chloride of gastrie juice is apparently derived ultimately from blood chloride (see Chapter 14), and is normally reabsorbed during later stages of digestion. Loss of gastric juice ebloride by vomiting, or in pyloric or duodenal obstruction, may lead to the development of nn nlkalosis due to bicarbonate excess, since the lost chloride is replaced in part at least by bicarbonate. It is interesting to note that on chloride-deficient diets there is no change in the output of chloride in the gastric juice, while obloride excretion by other channels, such as the urine and the perspiration. may be markedly deercased.

Another specific function of the chloride ion is in connection with the chloride shift in the blood during the carriage of carbon diovide (see Chapter 24) whereby the bicarbonate content of the blood plasma is significantly increased by exhange with plasma cbloride, which enters

the red cells.

Chloride is likewise concerned in osmotic pressure regulation, making up about two-tbirds of the total anions of blood plasma and occupying a similar position in the other extracellulm fluids of the body. The role of chloride in this connection, bowever, is subject to somewhat greater variation than the role of sodium, since to n certain extent the organism can exchange chloride ions for other anions within the framework of normal osmolar concentration of total anions. This ability to replace chloride ions by other ions is more restricted, however, than for the other major anion of the plasma, the bicarbonate ion, over half of which can be replaced, for example by the lactate ion under certain conditions.

The chloride ion is an activator of salivary amylase, but this action is not specific for chloride and may be due in part to the effect of strong electrolytes in general on the solubilities of such proteins as globulins. Radioactive chloride (Cl²⁸) has been used for studies on the volume of the

interstitial extracellular fluid of the body.

Chloride present in the diet in excess over that required by the body is excreted, chiefly by way of the kidneys Excreted chloride usually is accompanied by excess sodium or potassium unless there exists a need for conserving base, in which ease the ammonium ion accompanies the chloride ion to a variable extent. Factors controlling the excretion of chloride are similar to those concerned with the excretion of sodium and potassium. On a chloride-deficient diet, or in fasting, the chloride excretion may amount to but a trace. However the excretion by normal adults averages about 10 to 15 g, per day expressed as sodium chloride.

POTASSIUM

This element, like sodium and chlorine, is also present in plant and animal tissues entirely as the potassium ion. The adult human body contains about 250 g. potassium, present almost entirely in the cells rather tban in the extracellular fluids as with sodium. While normal human blood plasma contains only about 5 milliequivalents of potassium per liter of water, almost the entire cation content of red cells is made up of potassium (about 170 milliequivalents per liter of cell water). Thus potassium is the major "hase" of the body cells, and apparently subserves in the cells the same general functions relating to osmotic pressure regulation and acid-base balance that have already been described for sodium in the extracellular fluids of the body. This is not the only function of potassium, however; it has been shown to aid in the enzymatic transfer of phosphate from ATP to pyruvic acid, for example (see p. 994); and the tooic effects of a significantly elevated plasma potassium level (potassium poisoning) can hardly be ascribed to osmotic forces alone. The potassium ion has an effect on muscular irritability which, bke that of sodium, teads to antagonize the effect of the calcium ion. Nevertheless, under conditions of severe dietary salt restriction, calcium appears highly important in supporting the potassium content of tissue.5

The abundance of potassium in plant foods and in meat precludes the danger of a deficiency of this element in a mixed diet. From 1 to 3 g. is commonly excreted in the urine per day on such a diet. Growth in rate may be retarded by reducing the daily potassium allowance below 15 mg. in the male and 8 mg. in the female; this difference may be due to differences in growth rate. Potassium requirement varies considerably in the different animal species. Prolonged consumption of a potassium-deficient diet may result in failure of the animal to respond to a correction of the deficiency. Potassium deficiency in man is associated with weakness and muscular paralysis, accompanied by a fall in the plasma potassium level; in animals hypertrophy of the beart and kidneys has been noted. In Addison's disease the exerction of potassium falls and plasma potassium rises, with the reverse changes in urinary and plasma sodium, indicating a function of the adrenal cortex in this connection. The administration of sodium chloride alone will sometimes alleviate the symptoms of the disease, apparently permitting the organism to restore the altered relation hetween sodium and potassium retention and excretion. It has been claimed that the symptoms associated with removal of the adrenal cortex may be reproduced experimentally by measures which increase the plasma potassium level to about twice the normal value.

CALCIUM

Calcium is an essential constituent of all living cells. Its mode of action is not clear but it appears to play a part in decreasing the permeability of cell membranes and the irritability of cells in general. Its effect on neuronuscular mechanisms is shown in higher animals by the development of hyperirritability and tetany as a result of a decline in the calcium content of the blood. Such a fall is observed in tetany following removal of the parathyroids, symptoms becoming noticeable when the calcium-ion concentration falls below 3 mg. per 100 ml. (total calcium below 7 mg.).

⁵ Eppright and Smith. J Biol Chem., 118, 679 (1937)

Normal values are 9 to 11 mg, total or 4.25 to 5.25 mg, of ionized calcium. In fatal hyperparathyroidism, Ca++ values of 7 mg, (total calcium 16 mg.) have been noted. McLean and Hastings find the calcium of plasma nearly all accounted for as calcium ions and calcium bound with protein, an equilibrium existing between the two forms. The parathyroids appear of prime importance in regulating the calcium content of the blood. The mechanism of this action is not clear.

Vitamin D improves the utilization of calcium for calcification and other purposes. There is convincing evidence that vitamin D promotes calcium absorption; there is also some evidence that it may act on the calcification process itself. The mechanism of vitamin D actiou is obscure. Any relation between vitamin D and parathyroid action is, at most, indirect. The action of vitamin D in calcium metabolism is discussed further in Chapter 35. Since ealcium salts within the intestinal tract are more soluble at acid reaction than in neutral or alkaline solution, absorption of calcium is promoted by high gastric acidities, by acid-containing diets in general, and by diets containing lactose which lead to lactic acid fermentation in the intestines. Amino acid products of protein digestion may promote the absorption of calcium, possibly by the formation of soluble complexes. Other as yet unknown factors may also be concerned.

We are also much in the dark as to the nature of the calcification process as it occurs in hones and teeth. It appears to depend upon an adequate inorganic phosphate concentration of the blood plasma as well as upon the Ca++ of the plasma, and these are maintained through a balance between absorption from the intestines and excretion by the intestines and the kidneys. One view is that we have a simple ehemical or physicochemical equilibrium hetween blood and bone, the blood heing commonly supersaturated with respect to calcium phosphate and carbonate, with cartilage possessing a certain selective action. Another view is that calcification involves an active chemical process in the hone cells, such as the liberation of inorganic phosphate from organic phosphates through the action of phosphatase. The subject of calcification is discussed further in Chapter 9.

Calcium Requirement. The recommendation of the National Research Council is 0.8 g, per day for adults. There is a rather wide zone of calcium intakes over which different individuals can maintain or approximate equilibrium; adaptation to lower intakes has also been demonstrated. The question has even been raised whether or not the adult human male requires dictary calcium. Sherman found 16 per cent of American dictaries to fall below 0.45 g. The growing child requires considerably more calcium than the adult. Similarly increased amounts are needed by women during preguancy and lactation. Otherwise a negative balance for calcium may result and the drain upon the calcium of the bones of the mother becomes excessive Because of the large store of calcium in the bones the adult does not suffer so soon from calcium deprivation as the child. A lactating cow may lose as much as 20 per cent of its total supply of calcium without seriously interfering with milk

Illegsted, Moscoso, and Callaros. J. Nutration, 46, 181 (1952).

production In the growing child, however, any interference with the calcium supply leads to impaired calcification of the bones, as in rickets, as well as to slower growth of the body in general. It must be borne in mind that the child at birth is calcium-poor, i.e., the bones are incompletely calcified. The greater flexibility of the bones facilitates delivery, but the need of the newborn for calcium is necessarily more critical on this account.

Milk is of supreme importance in maintaining a proper calcium level ia the diet. Every child should receive along with other foods a quart of milk (containing about 11 g calcium) each day Milk is recommended in supplying calcium, because of its high content of calcium, the available form in which this is present (utilization about 30 per cent) and becau-e milk has otherwise a high nutritive value. Milk contains about 120 mg of calcium per 100 g, cottage cliecee contains about 100 mg Of other foods, eggs are useful, containing 70 mg per cent, mainly in the yolk, as are also green vegetables cabbage or lettuce supplying about 45 mg per cent of calcium Some green vegetables, like spinach, though high in calcium, are not so satisfactory, since much of their calcium exists as ovalate and is not readily utilized by the body. In certain foods, such as some cereals the presence of phytic acid (mositol hexaphosphorie acid) impedes calcium and magnesium absorption since this substance forms an insoluble compound with calcium and magnesium. This effect can be readily overcome by adding calcium carbonate to the diet, as for example by incorporation into bread Bread has about 30 mg per cent of calcium, and meat on the average about 10 The utilization of calcium is improved by vitamin D, an adequate supply of which should therefore be insured

The subject of calcium excretion has been discussed in Chapter 28 Most of the excreted calcium is found in the fees. This feed calcium presumably represents unabsorbed dietary calcium. There is an intestinal excretion of calcium which may amount to 0.5 g or more per day, mot if not all of this appears to be reabsorbed. The many factors which influence the absorption and excretion of calcium have led some to doubt the validity of calcium halance studies as an index to calcium requirement unless these various factors can be subjected to rigorous control.

The urmars output of calcium depends upon the nature of the diet but averages about 0 1 to 0.3 g per day for normal adults

PHOSPHORUS

Since the greater part of the pho-phorus of the hody is associated with calcium in bone the metabolism of these two elements is to a considerable extent parallel and follows that of the osseous tissues Pho-phorus however is abundant also in many of the softer tissues of the hody and plays many important roles in life processes. Through the intermediary formation of leethins it is concerned with fat metabolism. Through the formation of hexosepho-phates of adenvia and and of creatine phosphate, it plays a primary role in the carbohy drate metabolism of animals as well as in fermentation processes. Pho-phates play a role in the neutrality regulation of the organism (see p. 816). They are concerned with the ab orption of sugars from the intestine and the reabsorption of glucose in the

kidney tubules. Phosphorus is a constituent of the phospholipides present in all tissues and which are especially abundant in nervous tissues. It is present in nucleoproteins of the chromatin material of cells and in phosphoproteins such as casein. It has been suggested that most of the phosphorus, as well as the fat of milk, arises from the lecithin of the blood of the lactating animal.

Many studies of phosphorus metabolism have been made using radioactive phosphorus; this isotope also has some use in the treatment of disease (see p. 983). The important conclusions regarding phosphorus in its relation to bone and teeth formation which have resulted from the use of radioactive phosphorus have been presented in Chapter 9 For a further discussion of phosphorus in its relation to calcium metabolism and vita-

min D, see Chapter 35.

Phosphorus Requirement. The National Research Council states that the daily phosphorus allowance for children and for women during pregnancy and lactation should be fully equal to the calcium allowance. The allowance for other adults should be 50 per cent higher than the calcium allowance (see Table, p. 1108). Four per cent of American dietaries have been found to fall below the minimum phosphorus requirement, the danger of deficiency being less than in the case of calcium, and yet a matter of some concern. The phosphorus content of the diet is most readily maintained at a proper level through the liberal use of milk, a quart of milk a day containing 0.88 g. of phosphorus being recommended for children. In addition to milk (93 mg. of phosphorus per 100 g.) good sources of phosphorus are meat (average 175 mg. per cent), eggs (180 mg), ebeddar cheese (610 mg.), nuts (400 mg.), whole cereals (as whole wheat, 375 mg.), while white flour and polished rice are much lower in phosphorus (about 90 mg.). The phospborus in the bran of cereals (phytic acid) is utilized with some difficulty but, in general, the form of phosphorus in the diet is of little practical importance since in any case, as the result of the digestive processes, the phosphorus is absorbed in inorganic form. Too high a ratio of calcium to phosphorus in the diet, however, is unfavorable to phosphorus absorption. A diet high in beryllium or strontium also hinders phosphorus absorption and gives rise to a form of rickets.

MAGNESHIM

Magnesium is an indispensable constituent of all living cells. The daily requirement for magnesium has been estimated to be about 0.43 g for adults. Magnesium is present in such amount in plant foods and in meat that there is little possibility of the diet being inadequate with regard to this element.

About 71 per cent of the magnesium of the body is present in the boues. In muscle magnesium exceeds calcium in amount. In blood, where the reverse is true, the magnesium content is very constant. This is true also of the body as a whole, the concentration of magnesium remaining constant at about 0 045 per cent during growth, whereas calcium and phosphorus tend to increase in percentage. Excessive intake of magnesium gives rise to an increased excretion of calcium in the urine, and vice versa

Like calcium, magnessum depresses nervous irritability, but to a greater

extent A level of magnesium in the blood of 20 mg. per cent produces anesthe-ia, apparently by action on the central nervous system, whereas injections of calcium salts have a wakening effect. Magnesium ions are necessary for the action of many enzyme systems, particularly those concerned in carbohydrate and fat metabolism in muscle and other tissues The mode of action of the magnesium ion in these instances is not known. Magnesium ions inhibit the activity of adenosinetriphosphatase, the enzyme which splits phosphate from ATP. It has been suggested that this may be related to magnesium anesthesia.

Diets extremely low in magnesium (1.8 parts per million) have been fed to young rats; under these conditions serum magnesium drops to 0.5 mg per cent and there is marked vasodilation and hyperexcitability, leading to tetany and death. There is a rise of cholesterol of the blood and a corresponding decrease in free fatty acids, perhaps representing a disturbed metabolism of fat. There is also a marked decalcification of the skeleton which recalls the fact that grass tetany of cattle is associated

with a low magnesium content of the blood

Intestinal absorption of magnesium salts does not present a nutritional problem because of the relative solubility of the magnesium salts and their abundance in the diet The excretion of magnesium has been discussed in Chapter 28. The action of magnesium sulfate in the duodenum in hringing about emptying of the gallbladder is employed in clinical diagnosis and therapy.

IROX

Iron exists in the body chiefly in the heme portion of the hemoglobin of the red blood cells It is also found, however, in plasma in combination with serum globulin and as an indispensable constituent of various oxidation-reduction enzymes essential for the life of cells in general. Among the heme-containing enzymes are catalase, peroxidase, the cytochromes and cytochrome oxidase, and probably others. The action of these enzymes is inhibited by cyanide, which apparently combines with the iron. Iron is also found in the liver in the form of a compound called ferritin. Ferritin is an iron-containing protein, the protein portion of which has a molecular weight of about 500,000, and the iron is considered to be present in the form of colloidal Fe(OH), interspaced in the crystal lattice of the protein The full significance of ferritin in nutrition remains to be elucidated

Young animals (of all species so far studied) placed on a pure milk diet at the time of weaning develop a severe anemia. Addition of purified iron salts alone to the diet does not cure the condition, but does so if supplemented with very small amounts of copper salts. It is well estabhelied that the animal organism cannot convert the iron of the diet into hemoglobin unless very small amounts of copper are also present. There are a few instances in the literature where this has also been demonstrated in the case of human nutritional anemia. In the majority of human nutritional anemias, however, a copper deficiency does not exist and the anemia is due simply to iron deficiency. These nutritional anemias are not to be confused with the anemia of folic acid deficiency (see Chapter 35) or with permenous anemia or other anemias due to increased blood destruction or disturbances of the blood-forming organs

Iron Requirement. The recommended daily allowance of iron as given by the National Research Council is 12 mg. per day for adults (15 mg, per day in pregnancy), and 6 to 15 mg per day for children, depending upon age. It has been estimated that for infants 0.5 mg. per kg. body weight per day is necessary to insure retention of iron. The form of the iron in the dict must also be considered. Anemia due to iron deficiency is not uncommon in women and preschool children. Hence general adequacy of this element in the diet cannot be assumed. Infants are born with a high level of hemoglobin iron and with an additional iron reserve in their livers which is later used for hemoglobin formation to supplement the low iron content of their milk diet. If continued beyond the normal lactation period such a dict leads to the development of a untritional anemia. Children ordinarily require supplementary iron-containing foods by the time they are one year old. There is some evidence that the administration of iron to normal children and young adults, or increasing the iron intake above the usual levels by other means, will consistently produce a slight but significant (about 10 per cent) increase in the blood hemoglobin content.

Milk contains from 0 07 to 0.1 mg. iron per 100 g. This small amount of iron appears to exist in a readily assimilable form. Most foods appear to contain two forms of iron; home iron which is not utilizable and nonheme forms which are available. Metallie iron or any of the ordinary iron salts may be used to supply iron. It has been shown that under certain conditions ferrous compounds are somewhat better utilized than ferric compounds. Eggs contain 3 mg, iron per 100 g, of food, all of the iron being available. Lean muscle meats contain about the same amount, 50 per cent of which is available. Beef liver or heart coatain about 8 mg. per cent of iron, about 60 per cent of which is available. Spinach contains about 3 6 mg. iron but like the iron of blood, alfalfa, and oysters this appears to be less than 25 per cent available. The minimum amount of iron which must be added to milk for maximum hemoglobin regeneration is said to be 0 0007 per cent.

The absorption and excretion of iron have been the object of much study. Relatively little is known about the mechanism of iron absorption, but it is known for example that the action of copper in promoting iron utilization is not related to iron absorption. There is some indication that the absorption of iron may be influenced by the state of the body iron stores Studies with radioactive iron have shown that iron may be absorbed very rapidly, appearing in the red blood cells in about four hours. However it takes about a weck for the complete conversion of absorbed iron into hemoglobin.

Very little iron is ever excreted Only 1 to 3 mg. per day is found in the urine, with more in the feces, but this latter probably represents for the most part unabsorbed iron of the diet. Iron split off from the hemoglobin of the red cells after their destruction appears to be retained within the body and used over again. The chief need for iron appears to arise in infants on a milk diet, or in others after blood loss as by hemorrhage or during menstruction.

Moore, Dubach, Minnich, and Roberts: J. Chn. Inrest , 23, 755 (1944).

COPPER

Although copper has long been known as a constituent of such compounds as the hemocyanin of the blood of certain lower organisms the first demonstration that copper played a vital part in the animal organism was in connection with the conversion of dietary iron to hemoglobin as described on p 1002. Since that time copper has been shown to be an essential component of certain enzymes such as tyrosinase and accorbic acid oxidase and a copper containing protein (hemocuprein) has been soloated from animal blood. Animals placed on a copper-deficient diet soon lose weight and die but death is not due to the concomitant anemia since an equally intense anemia due to simple iron deficiency may be maintained for a long time. Thus copper is an essential element in the animal hody but the precise role still remains obscure "Some studies have demonstrated that copper is associated with the activity of certain oxida ton reduction enzymes in tissues.

The amounts of copper required per day are extremely small, for an adult only about 1 to 2 mg whereas a child requires only 0 00 mg per kg of hody weight The adult human body contains about 0 1g of copper Copper appears generally to 1e present in the ordinary mixed diet in adequate amounts so that the possibility of a copper deficiency is limited Perhaps because of this fact there have been relatively few demonstrations of copper deficiency in man

IODINE

The adult human body contains about 20 mg todine about 10 mg of which is in the thyroid The daily allowance recommended by the la tional Research Council is about 0 003 mg per kg body weight It is needed for the production of thyroxine and a deficiency leads to the condition of simple gotter Ocean water is relatively rich in todine as is also the dust formed in the atmosphere from the drying of the salt spray which dust is carried inland by the winds where the rains dissolve it and carry it into the soil which thus becomes enriched with rodine Where the distances from the sea are greater or mountain barriers intervene this does not occur at the same time rodine is being leached out of the soil which thus becomes low in iodine. In such regions, including the Great Lakes region and the Upine regions of Furope the vegetable foods grown in the iodine poor soil are poor in iodine as is also the drinking water and gotter is prevalent In Japan gotter is very rare in part because of the use in the dict of seaweed which is very high in todine Sea foods such as f sh and ossters are rich in iodine Good sources are al 0 vegetal les from sea coast districts Fair amounts are also found in the fat of milk In gotter regions however such foods should not be depended upon but should be supplemented with potassium iodide. A few drops of 10 per cent todide solution given every two weeks would cover the requirement since some storage of ingested rodine occurs. The most

^{*(} lass Copper Metololum Balt more Johns Hopkins University Press 19.0 Marston I hastel Ress 12 6f (19.7)

satisfactory method for insuring an adequate intake of iodine without danger of overdosage is to use table salt to which has been added one part per 100,000 of iodine as sodium iodide. Such salt is widely sold. Care must be taken in the administration of iodine to persons with hyperactive thyroids Studies with radioactive iodine indicate that this element is rapidly taken up from the blood by the thyroid gland, with significant differences in this respect between normal, hyperthyroid, and hypothyroid individuals (see also p. 983). Iodides also appear to be selectively concentrated and excreted by the salivary glands in man.

Variations in the iodine content of the blood show a relation to thyroid disease, but the results obtained are somewhat irregular. The normal range of serum iodine content is from 5 to 12 µg, per 100 ml. About 85 per cent of the serum iodine is bound to proteins, chiefly albumin. This constitutes the protein-bound iodine of the serum, the concentration of which is of diagnostic value regarding the level of thyroid function. The normal level of serum protein-bound iodine is from 4 to 8 µg, per cent. A thyroid hyperplasia has been produced in rabbits by feeding cabbage. Various synthetic compounds are known, such as thiouracil and thiourea (see pp. 771–2) which evert a specific depressing effect on thyroid secretion and have proved useful in the treatment of thyroid disease.

SILFUR

Strictly speaking the metabolism of sulfur does not come under the bead of inorganic metabolism, since only an insignificant part of the sulfur ingested is in inorganic form, by far the greatest proportion being combined in protein molecules as the amino acids cystine and methonine. Proteins of foods vary in sulfur content from 0 4 to 1.6 per cent, with an average of about 1 per cent. The sulfur-containing amino acids and their metabolism are discussed in Chapter 33.

The end product of sulfur metabolism is taurine or sulfuric acid; the latter is either immediately neutralized and excreted as inorganic sulfate in the urine or may be first conjugated with phenol, glucuronic acid, or indovyl. On the average, about 1.0 g. sulfur is excreted daily in the urine.

Certain sulfur compounds have important hiological interest, e.g., thiocyanate in saliva and other fluids, taurocholic acid in bile, ergothioneine of the hlood corpuseles, and glutathione present in all cells and concerned with oxidation processes These substances are discussed elsewhere in this hook, as are also the properties and metabolism of the sulfurcontaining amino acids and the various forms of urinary sulfur.

MANGANESE

There is considerable evidence that manganese in small amounts is a dietary essential. Rats placed on a manganese-deficient diet show a retardation of growth and reproductive failure in both male and female. The arginase activity of the liver of the manganese-deficient rat is lower than normal; in ratro addition of manganese ions restores the enzymatic ac-

^{&#}x27; Salter, Bassett, and Sappington. Am J. Med Sci . 202, 527 (1941).

trity. Other isolated enzyme systems are known to be netwrited by manganese ions in small amount. In the case of the growing clink a manganese deficiency is associated with the development of percoss (bone malformation). Studies with radioactive manganese indicate that the chief channel of excretion is through the liver into the bile. It has been suggested that the daily diet of children should provide from 0.2 to 0.3 mg manganese per kg of hody weight.

OTHER LILMENTS

Fluorine Fluorine is found widely distributed in animal and plant tissues, in very small concentration When added to the thet in small amount it is excreted in proportion to the intake Individuals on a normal diet containing no added fluorine excrete about 0 3 to 0 5 mg per day, this presumably represents the amount ordinarily present in the diet In certain parts of the world the soil contains sufficient soluble fluoride so that the drinking water, and food grown on the soil contain enhanced amounts of fluorine relative to other localities Individuals living in such communities are prone to develop "mottled enamel" (deutal fluorosis), an unsightly condition of the teeth. At the same time, the incidence of dental caries in these communities is in general much lower than elsewhere The concentration of fluorine in water claimed to be effective in the prevention of caries approximately 1 p p m is said to be below the level which will cause dental fluorosis \ fluorine level of 6 p p m appar ently has no harmful effects on the bones. The fluoride ion in significant concentration (0.01 M) is an enzy matte poison for tissues and fluoride salts are used commercially as ingredients of animal poisons (For a further discussion of fluorine see Chapter 9)

Cobalt Cobalt is present in plant and animal tissues in small amount, and there is conclusive evidence that it is an essential element in animal mutrition. It is present in vitamin Bi₁ (see p. 1209). Certain diseases of cattle and sheep have been attributed to a cobalt deficiency, and respond to cohalt therapy. In cattle cobalt dietary deficiency is accompanied by a microcytic hypochromic anemia rapid loss of weight absence of estrus abortion, and low milk yield. In the case of sheep disease the cobalt is effective orally but not when mjeeted. It has been shown that cobalt increases the synthesis of vitamins B₄ and B₁₂ in the runeu of sheep. The administration of cobalt salts to rats and rabbits produces a marked polycythemia the reason is not known. Studies with radioactive cobalt indicate that absorption and chimination take place rapidly, with the kidney the main channel of excretion.

Zinc There is definite evidence that zinc is an essential element in animal nutrition. Rats on a diet low in zinc show a marked delay in intestinal absorption and a retarded growth rate. Zinc is a constituent of highly purified carbonic anhydrase the enzyme important in the formation and decomposition of carbonic and set be enzyme part of the zinc content of carbonic anhydrase is about 0 2 to 0 3 per cent 1 e. very similar to the iron content of carbonic anhydrase is about 0 2 to 0 3 per cent 1 e. very similar to the iron content of himoglobin (0 34 per cent). Zinc is also a constituent of crystalline (but not of amorphous) insulin. Whether zinc plays any significant role in connection with the action of insulin or carbonic anhydrase.

is not known, nor has any specific function been ascribed to zinc in animal nutrition. However, zinc is said to enhance the action of the follicle-stimulating hormone (FSH) and of the luteinizing hormone (LH) of the anterior pituitary.

Trace Elements. Various other elements found in traces in biological material are of uncertain significance. Aluminum has a wide distribution in animals and plants but is not known to be essential, nor does any aluminum added to food during cooking in aluminum utensils have any demonstrable effect on nutrition. However, it has been demonstrated that female albino rats fed bisenits made with a tartrate baking powder gave evidence of a distinct nutritional advantage over rats from the same litter which were fed biscuits made with an alum baking powder. This advantage was apparent from the standpoint of reproduction as well as from that of growth.

Boron is essential for plants but apparently has no significance in animal nutrition. The position of arsenic as a trace element is debatable. Silicon is found in plant and animal tissues but its function is unknown. Bromine has no normal or pathological significance aside from its known pharmacological effects. Selenium may be taken up by plants and thus become transferred to animals where it has a toxic effect which is counteracted by methionine. Molybdenum is an essential component of the

enzyme xanthine oxidase.

Acid-forming and Base-forming Foods. Certain foods, such as vegetables and fruits, on burning outside or inside the body leave an ash or residue in which the basic elements (sodium, potassium, calcium, and magnesium) predominate; whereas cereals, meat, and fish foods leave an ash in which the acid-forming elements (chlorine, phosphorus, and sulfur) predominate. Such foods are spoken of as base-forming and acid-forming foods, respectively, and will influence the acid-base balance of the body and the acidity of the urinc. Sulfur, while present in foods ebicfly in neutral form in the sulfurized amino acids, is oxidized in the body to yield sulfuric acid and is hence an acid-forming element. High-protein foods are hence generally acid-forming. On the other hand, the citrus fruits contain citric acid and acid potassium citrate, the citrate radicals of which are completely metabolized in the body like carbohydrate, leaving behind potassium, which is one of the bases of the body (see p 1084). Hence many "acid" fruits are base-forming Grape juice is much less effective than orange juice in reducing urine acidity because the tartaric acid it contains is not completely oxidized but is eliminated to a certain extent in the urine as such Pruncs, plums, and cranberries contain quinic acid. This is excreted in the urme chiefly in the form of hippuric acid (a conjugate of benzoic acid and glycine) which increases the acidity of the urine. With the exception of foods containing incompletely oxidizable organic acids, the acid-forming or base-forming value of foods may be calculated by obtaining the differences between the equivalents of normal acid, calculated from the content of sulfur, chlorine, and phosphorus (considering

¹⁶ Hawk, et al. Unpublished For abstract see Philip B. Hawk and Collaborators: Researches and Writings Published privately, 1942.

phosphoric acid as divalent) and of normal alkali calculated from the content of sodium, potassium, calcium, and magnesium. A table of the acid- or base-forming value of various foods will be found in Appendix IV.

Through the use of considerable amounts of potatoes or other vegetables or of fruits such as oranges, it is possible to lower the acidity of the urine markedly or even to make it alkalme. Naturally reduction in the amount of acid-forming foods has a similar tendency. Increase of urinary acidity above the usual levels is less readily brought about. Reduction in the acidity of the urine by increasing the solubility of uric acid therein may reduce a tendency to formation of uric acid cleruin foods may be otherwise desirable, since too much acid-forming food might under certain circumstances be a drain upon the fixed base of the body. Fortunately, however, the body has ordinarily a marked ability to protect itself from excess acid formation, through ammonia production and in other ways It is not clear, therefore, that a harmful effect is produced by a preponderance of acid-forming foods provided minerat, vitamin, and other dietary requirements are met

EXPERIMENTS ON INORGANIC METABOLISM

 Solt-free Dict. The effect of a salt-deficient dict upon the metabolic processes is reflected in the composition of the urine as shown by the following experiment

Procedure. Ingest an ordinary mixed diet containing an ample sait content for a period of two days. Follow this period by the Ingestion of a diet which has had its sait content reduced to a very low value. Sugar and olive oil or nonsaited butter may supply the buik of the calorific part of the diet and dialyzed egg white or casein or commercial protein preparations, e.g., plasmon, gluten, or glidme may supply the protein. Ingest such a diet for three days. (This is an acid-forming diet, p. 1977.) Collect the urine and analyze for chloride, thratable acidity, ammonna, and total nitrogen. Compare the data from the normal days with those obtained when the sait-free diet was incested.

2. Salt-nch Diet. On an ordinary mixed diet a normal adult will daily excrete 10-15 g of chloride, expressed as sodium chloride, in the tunne On a salt-free diet this exerction decreases, whereas if the diet contains an excessive quantity of sodium chloride this excess will be promptly excreted in the tunne Normal feces contain very little sodium chloride even after excessive sodium chloride ingestion (see Exp 3, below)

Experiment. Ingest an ordinary mixed diet for two days. On each of the following two days take a similar diet plus a welghed amount (e.g., 10 g.) of sodium chloride in divided doses. Collect the urine for the four days in 42thr. samples, preserve and analyze for sodium chloride (for methods see p. 955). What proportion of the added chloride was recovered?

3. Inorganic Elements in the Feces. The salts of sodium and potassium are almost completely absorbed from the intestine I lence the alkali metals and chlorides are excreted mainly in the urne and are found only in very small amounts in the feces even when large amounts are ingested With calcium, magnesium, iron, and phosphate, conditions are different. Ordinarily about 90 per cent of ingested calcium and

over half of the magnesum are climinated by way of the feces. From 20 to 30 per cent of the phosphorus ingested is usually found in the feces.

Experiments (a) Ingest for a period of three days on ordinor; mixed diet without odded solt ond containing no milk. Seporote the feces for the period (see p. 1046) ond retoin o portion of the well-mixed feces for onolysis

(b) Proceed os above with the exception that there is added to the mixed diet 10 g of common sait and a quort of milk (containing obout 11 g Ca, 01 g Vg, 14 g chloride expressed as sodium chloride, ond 10 g P) Mix the feces well ond reserve part for analysis

Ash 10 g samples of the feces from the above diets Dissolve with the ald of a little dilute nitric acid, filter, and make up to 100 ml Determine in oliquot portions of this solution (1) Chlorides (2) Colcium and magnesium (3) Phosphorus (For details of analytical methods see Chapter 31) Calculote the percentages of the odded Ca, Mg, P, and Cl which are recovered from the feces

For a more detailed study of chloride excretion combine this experiment and Exp 2, above

4 "Alkoline Tide" For a time after a meal the normal acid reaction of the urine may be changed to neutral or alkaline. This has been explained as due to the production of a tempority alkalosis because of the secretion of the acid gastric june. The presence of no alkaline tide has been used as an indication of the secretion of hydrochloric acid in the stomach to cases where it was desirable to avoid passing the stomach tube. The uniony ammonia excretion may also serve a similar purpose

Experiment Ingest an ordanory mixed diet Urinate just before dinner and note the recotton of the urine to litmus If acid, determine the hydrogen ion concentration by the method given on p 871 (If alkaline, discord the urine and make the test on another day) After coting a heavy dinner (meats) collect the urine at intervals of holf an hour and toke the reaction to litmus and determine the hydrogen-ion concentration os before Did your urine change in reaction after the meal and if so how long a period elapsed between the meal and the occurrence of the maximum change in reaction?

5 Hydrogen-ion Concentration of the Urine as Influenced by the Ingestion of Acid forming and Bose farming Foods. It has been demonstrated that vegetables and fruits, on burning leave an ash io which the basic elements (sodium, potas sum, calcium, and magnesium) predominate whereas cereals meats, and fish foods leave an ash in which the acid forming elements (chlorine, sulfur, and phosphorus) predominate.

The acid or base-forming potentialities of various foods are given in the Appen dix Potatoes, oranges, raisms, apples bananas, and cantiloupes are important baseforming foods. Among the most important acid forming foods are found rice, wholewheat bread, outmeil meats, and eggs. Certain fruits, e.g., cranbernes, primes and plums, yield a base ash but are acid forming foods. For further discussion see p. 1097

On a mixed diet the hydrogen ion coocentration of the urine has been found to average about pH 60 Base-forming foods decrease the acidity and the ammonia content of the urine, while and forming foods have the reverse effect

Experiment Ingest a uniform diet consisting of milk, crackers, butter, peanut butter, and water in desired quantities for a period of three days Follow this by a period of six days during the first three of which considerable quantities of acid-forming foods (see Appendix) are added to the diet, During the second half of the period (days four to six) add an abundance of base forming foods to the diet Distilled water should be used for drinking purposes and a uniform volume should be ingested daily Collect the urine in 24 hour periods preserve, and analyze for hydrogen ion concentration, titratable acidity, and ammonia (for methods see Chapter 31) Compare your results with those tabulated in the table below

REACTION OF URINE AS INFLUENCED BY DIETII

	Basa Dut		1	2	3	4	5	6
				Barol Dut I		Basal Diei No 2 Plus		
Determs nation	5 days	5 days	Baked potatoes (7:0 g. per day) (6 days)	R ce (210 g per day) (4 days)	Cranberry sauce (300-600 g. per day) (6 days)	Bread 4 (whole wheat) 450 g (t day)	Prunes (330-5.0 g per day) (3 days)	Cautalous (260 s per day) (5 days)
pН	7 19	5 57	7 74	7 48-6 90 7 14	6 30-5 70	6 80 (Prev ous day 6 90)	5 30-4 50	5 30-7 3
T tratal l acd ty (mL 01 N	275	474	196-216	166-297	391-488	350 (Previous day 297)	570-540-578 563	328
Am non N (gram	a) 0 310			0 166-0 25 0 198	0 219-0 39	0 280 (Previous day 0 251)	0 602-0 725	0 513-0

⁶ Hydrogen ion Concentration of the Urine or Influenced by Alkali Ingestion. The ingestion of certain organic salts of the alkalies e.g., sodium citrate and sodium blearl onsite causes an increase in the pH of the urine. The ingestion of acids (either organic or inorganic) or acid salts e.g. sodium dihydrogen phosphate decreases the pH of the urine. The alkalies are much more effective in producing changes in rection than are the acids. The influence of ingested alkali (sodium bicarbonatte) is shown in the table on p. 1101 containing data submitted by Henderson and Palmer.

snown in the table on p 1101 containing data submitted by Henderson and Psinet Blatherwick reported a decrease in ammonia nitrogen output from 0 256 g to 0 072 g and secompanying decreased acidity under the influence of bicarbonate ingestion (25 g in two days)

INFLUENCE OF INCESTED SODIUM BICARBONATE ON HYDROGEN-ION CONCENTRATION OF URINE

	Sodium		Time of Collection of Specimen of Urine and pl						
Experiment Number	Bicar- bonale, Grams	Bicar- bonate Ingestion	11 00 AM	12 00 Noon	1 00 PM	200 PM	3 00 P M.		
1	4	7 40	8 30	7 48	7 48	7 40	5 85		
2	8	5 40	8 50	8 30	6 50	6 50	7 40		
3	12	5 30	8 70	8 70	8 70	8 70	8 70		
4	8	7 40	8 50	8 70	8 50	8 50	8 50		
5	8	5 85		}	8 70	8 70	8 30		
6	8	6 70	7 48	8 70	8 50	8 70	8 50		

periods and analyze it for titratable acidity, hydrogen-ion concentration, and ammonia Compare your results with those shown in table above

If desired, the bicarbonate may be given in one dose of 8 to 12 g. and the urine collected in hourly specimens for the next 5 hours and each specimen analyzed. Data from such experiments are shown in table above

 Influence of Calcium Deficiency. A demonstration of the harmful effect following the elimination of calcium from the diet may readily be made if the diets listed belon be fed to young white rats

Procedure. Place two young white rats (40 60 g.) in separate cages and feed the diets given in the following table. Make frequent body-weight determinations. The rat receiving Diet 2 will show normal growth. The rat receiving Diet 1 will fail to show normal gains in weight. This diet is deficient in calcium. See Figs. 269 and 270

	Diet 1	Diel 2
	Per Cent	Per Cent
Beef liver (steamed and dried)	20 0	20 0
Casein	100	10 0
NaCl	10	10
KCi	10	10
CaCO ₁	0.0	15
Dextrin or starch	65.0	63 5
Butter fat	3.0	3 0

8 Influence of Ultraviolet Radiation on Inorganic Metabolism. Ultraviolet radiation has calcium-depositing properties. See p. 1248

Procedure Place three young white rats (litter mates) weighing 50-60 g, in individual cages, and supply ad lib a rachitogenic diet such as described in Chapter 35. Keep the rats in a dark or dimly lit room, but subject one to an hour's exposure to direct sunlight at about noon each day, and expose another to I minute's irradiation at a distance of 2 feet from an ultraviolet lamp After 4 weeks examine the leg bones roentgenographically or by the line test (see p. 1265), and perform bone ash nnalyses on the dried, fat-free bones,

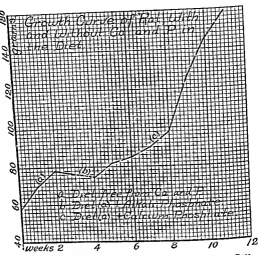


FIG 269 GROWTH CLEVE OF RAT WITH AND WITHOUT CALCILM AND PHOSPHORES IN THE DIET

Unput hat ed data from the senior author's laboratory



Fig. 270 SKELPTONS OF TWIN ALBINO RATS SHOWING INFLUENCE OF CALCULAR DEFICIENCY
COUNTRY Florings and Minchest J. Biol. Chem., 44, 429 (1925)

9. Influence of Iron and Copper Deficiency.18 The anemia caused by the ingrestion of milk as the sole article of diet may be cured by the addition of an iron (and copper) supplement

Procedure. Pince three young white rats (litter mates) on a milk diet and perform hemoglobin determinations weekly on blood obtained by clipping off the tips of the tails. After the hemoglobin has dropped to half its original value, feed one rat a daily supplement of 0 25 mg. of Fe (as ferric chloride) and another rat 0.25 mg. of Fe (as ferric chloride) and 0.05 mg. of Cu (as copper aulfate). Continue the third animal on the original milk diet without Fe or Cu supplement. Continue weekly determinations and compare the condition of the three animals, noting especially the color of the tails, ears, and eves.

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¹² For quantitative technique for the study of nutritional anemia see Elvehlem and Kemmerer J Biol Chem , 93, 189 (1931), Smith and Otis J Autrition, 13 573 (19:

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Vitamins and Deficiency Diseases'

In 1881 Lunin, a pupil of Bunge, recognized that a diet consisting only of purified protein, earbohydrate, fat, and minerals would not sustain life When such diets were supplemented with certain natural foods, notably milk, normal growth and well-being of the experimental animals were restored These findings were in accord with earlier demonstrations of the curative value of citrus fruit juices in scurvy, a disease contracted by sail ors, soldiers prisoners, and others who subsisted on limited fare They also coincided with observations by the Dutch physician Eijkmann in 1897 that beribers or polyneuritis, a disease resulting from the prolonged consumption of polished rice, could be cured by supplementing the diet with rice polishings or extracts thereof. These and similar studies suggesting the presence of indispensable substances in natural foods were brought to a head by Hopkins (1912) who called them "accessory food factors" The work of Osborne and Mendel and of McCollum (1913) established beyond doubt the presence of these factors in milk Funk (1912) introduced the term vilamine in the behef that he had isolated one of these factors and established that it possessed the properties of an amine The final e has since been dropped masmuch as these substances as a class are not related to ammes

A vitamin is a potent organic compound, occurring in minute proportions in natural foods, which must be available to the animal organism from the diet or other sources in order that a apeense metabolic process or reaction may proceed normally. The "other sources" referred to in this definition are, for example, ultraviolet irradiation of precursors of vitamin D in the skin or bacterial synthesis of certain vitamins of the B group in the intestinal tract. Unlike amino acids vitamins do not enter into the tissue structure. Vitamins are exogenous, hormones endogenous, Compounds which may be vitamins for some apecies may be synthesized by others and hence behave in effect as hormones.

With the rapid advancement of our knowledge of the chemical constitution of the vitamins, it is not surprising that structurally related chemical derivatives, both natural and synthetic, should be discovered which exhibit typical physiological activity, though perhaps in varying degree For these, the term relamers has been suggested Naturally occurring

¹Tl e ra; id strides being made in this field require that special attention be pull to the date of pull iriation of insternal on the sulject. Important books an I review articles on the general subject of vitamina are cited in the Bill argraphy at the end of this of appear.

vitamin D exists in various chemical forms, and what was previously regarded as vitamin A in plant sources is now known to consist of various carotenoids, principally \$\beta\$-carotene, which function as precursors of the vitamin Hence a distinction should be made between vitamin activity, which indicates certain specific biological properties, and the vitamins per se, which refer to definite ebemical entities, of which there may be see each exhibiting similar hological activity.

Some vitamins are characterized by species specificity, i.e., they are required by certain species of animals and not by others. For example, vitamin C is necessary in the diet of man, guinea pigs, and monkeys, but dogs, rats, and other species are able to synthesize it. Hence for these species vitamin C may be regarded as a hormone. To a certain extent the physiological requirement of various species of animals for vitamins may be met by bacterial synthesis in the rumen or intestinal tract. This process, however, does not signify that these species do not require the nutrient in question.

The vitamins are recognized biologically by their absence rather than by their presence, that is, the characteristic symptoms of each vitamin deficiency must be produced in order that the effect of the vitamin in question can be demonstrated. In addition to (and probably secondary to) their role in the prevention of specific deficiency symptoms, each of the vitamins participates to a greater or less degree in the promotion of growth. Our knowledge of the physiological modus operands of the known vitamins is not far advanced. It is known, however, that most of the vitamins participate in enzyme reactions of a highly complex nature. Vitamins may be precursors of enzymes, as thamine is of cocarboxylase or maem of coenzymes I and II, or they may be substrates, as vitamin A is of rhodopsin.

The condition induced by the absence of a single vitamin from the diet is called an aritaminosis, e.g., avitaminosis B₁, if the deficiency is multiple, the term polyaritaminosis is applied. Many observations, particularly on the effect of deficiency or dosage of one vitamin on the storage or exerction of another, suggest the need for balanced proportions of certain dietary essentials. Until more is known about the specific mechanism of the action of individual vitamins, however, it will be difficult to establish the significance of such interrelationships.

The present list of vitamins cannot be regarded as complete, but it is probable that from the standpoint of human nutrition all the major vitamins have been identified. The table on p. 1106 summarizes the common names of the vitamins, and their principle vitamers—1 e, structurally related compounds possessing qualitatively similar activity. Some question has been raised as to the justification for including choline or mositol among vitamins, since such compounds are present in the foods in more than trace quantities. Nevertheless, they do not form a material part of body tissue, and functionally they conform to the definition of vitamins stated above.

Concurrent with the advances in isolation and identification of the vitamins, chemical and physical analytical procedures have been evolved for determining the content of almost every vitamin More recently.

SUMMARY OF PRINCIPAL COMPOUNDS WHICH POSSESS VITAMIN ACTIVITY

Vit	amın	Principal Vitamers				
Common Name	Principal Synonyms	Natural	Synthetic			
itamin A	Axerophthol Antiophthalmic fac- tor Vitamin Ai	u-, p-, and ,	Vitamin A acid Vitamin A ketone Vitamin A esters			
Thiamine	Vitamin B ₁ Ancurin Antineuritic factor Antiberiberi factor		Vitamin B. disulfide Analog with methyl group in position 6 instead of 2 in p) rimidine ring			
Ribnflavin	Vitamin B ₂ Vitamin G Lacto-, ovo-, verdo or hepatoflavin	Riboflavin mononu- eleotide -, Riboflavin dinucleo tide	7-Methyl 0 (p-1'- ribityl)-isoalloxa zine 6-Methyl-9-(p-1'- ribityl)-isoalloxa zine 6-Ethyl-7 methyl 9- (p-1' ribityl) isoal loxazine			
Nisein	P-P factor Antipellagra fact Anti black tongu		Coramine Esters of macin			
Vitamin B.	Pyridoxine Anti acroslynia l tor Adermin	Pyridoxal Pyridoxamine Pyridoxal phosphate	,			
Pantothenic nest	Chick antiderm tis factor Filtrate factor	stı	Esters			
Biotin	Skin factor Inti-egg white jury factor Bios II	D thohotin	Sulfoxide of hiotin Laters Dethiobiotin			
Pteroxighat acad	Anti-anemia la	Fermentation I, ca actor factor I ner I case facto	İ			

SUMMARY OF PRINCIPAL COMPOUNDS WHICH POSSESS VITAMIN ACTIVITY -(Continued)

V	ilamin	Principa	l Vitamers
Common Name	Principal Synonyms	Natural	Synthetic
p-Aminobenzoic acid	Chromotrichia fac- tor Vitamin B _x Anti gray-hair factor		
Vitamin Bi+	Cobalamin Cyanocobalamin	Vitamin B _{12s} (= B _{1*b}) = Hydroxocobalamin	
Choline	Sinkalin Bilineurine Fagin Amanitin	Methionine + eth- anolamine Betaine + ethanol- amine	Analog containing phosphorus instead of nitrogen Arsenocholine Methyl-diethyl hom olog Triethyl homologs
Inositol	Bios I Mouseanti aloperia factor Rat antispectacled eye factor	Phytin Soybean cephahn	Methyl mositol Inositol hexa acetate
Ascorbic acid	Vitamin C Antiscorbutic vita min	Dehydroascorbic acid	6-Deoxy ascorbic acid Isoascorbic acid I-Fucoascorbic acid
Vitamin D	Antirachitic vita-	Vitamin D ₂ (Calci ferol) Vitamin D ₁	Viosterol (irradiated ergosterol) Irradiated 7-dehydro cholesterol Irradiated 22-dihydro ergosterol (D ₄) Irradiated 7-dehydro- sitosterol (D ₅)
∞-Tocopherol	Vitamin E Antisterility vita min Fertility vitamin	β and γ Tocopherols Esters	Esters Analogs with ethyl substituents in place of methyl Analog with NH2 in place of OH
Vitamin K	Vitamin K ₁ Phylloquinone Antihemorrhagie vi tamin Coagulation vita min	Vitamın Kı	Menadione Menadione sodium bi- sulfite Esters of the hydro- quinone forms

Recourt top Dair Differ Lilonalers

DYYON	Vita-	I D		90	90	2 <u>3</u> 5	\$8 <u>8</u>	999	999
MATE OF C	lscor- bic lci I	вш	555	2223	120	228	800	2880	588
ARIFD DU	Уласия	Bu	222	2225	55	400	ాంల	222	3 23
VED IN V	Rebo-	Бш	000			500	220	-0101 0100	4614 800
CONSU.	Thia- mine	đu,	9-6	1111	240	-000 -100	980	111	
US WHF' donous AA A Council	Vita- min 1	ın	2000 2000 2000	2000	2000	1500	2000 2500 3500	2000 2000 2000	1500 5000 5000
n tiif l Lir \ 10 Researc	Iron	Вш	ដូងដ	222	55	909	1087	222	255
HETCHAIN OF GOOD NUTRINGS OF HOLDER PROGRESS WITH US WHEY CONSULED IN VARIED DIFFS OF CONTON DIFFICACION TO GOOD NUTRITION OF HOLDER PROGRESS WITH US WHEY CONSULED TO VEFT OF THE WAS NORMALLY TROODERS AND HITHOUT DIFFS OF CONTON FROM THE	Cal.	6	888	000		0000	000		222
	Protein	0	838	ន្តន្តន	88	N. X 33 53	223	ខននិ	525
UTRITION OF 1H OT 1791 Y TO A an I Autrition	Calories		3200 2800 2100	2300	3300	**** *** *** *** ***	1200 1600 2000	2500 3200 3800	2300 2500 2100
Good No	Height	(m) (m)	(170 (07)	(50) 251		855 988 588	87 193 193 193 193 193 193 193 193 193 193	141 (57) 163 (01) 175 (63)	111 (57) 110 (53) 112 (61)
UNTERANCE OF	Il eight	19 (10)	65 (143)	55 (121)	Pregiont (3ed Tennester)	6 (23) (23) (23)	21 22 25 25 25 25 35 26 35	75 (78) 44 (108) 63 (139)	76 (79) 17 (108) 54 (120)
Properties Al-	190	1	ยะอ	855	Pregnant (3c	1/12 3/13 4/12 9/13 10/12 1	-00	10 12 13-15 16-20	10-12 13 15 16-20
Divite	-		Mrs	Nomes		Infants*	Chil Iren	1803.	Curls

Numerors Nor Tanu arre (excerpted from the original)

ed While a requrement for certain unsaturated fatty needs (the inolese and arachidonic needs of natural fatis) has been amply demonstrated experimental animals the human need for these fatty acids is not known

cluded in the diet to the extent of at least 20 to 25 per cent of the total enlocres and (6) that the fat intake include essential unsaturated fatty neids to the extent of nt least 1 per cent of the point enlories. At higher levels of energy expenditure or for a very netive person consuming 4500 enlories and At work or in hot weather requirements may reach 5 to for eliblen and adolescent persons it is desirable that 30 to 35 per cent of the total calories be derived from fat Hater An ordinary standard for diverse persons is 1 ml for each calorie of food

The requirement for sodium chloride is increased by enviconmental and chimatic conditions associated with increased sweeting. Under unusual conditions such as doing heavy work in a hot elmate 10 to 15 g daly or even more may be required with meals and in deniking water However after neelmatination to lient the sodium content of sweat is greatly reduced and the allowance for sait can be near to normal Phenylous 13 liters daily Water should be allowed ad thatum mines sensations of thirst usually serve as adequate guides to intake except for infunts and sick persons Sodum Chloride Average intake of sodium chloride (salt) for the normal adult is 7 to 15 g daily

pregame, and during lactation. For other adults the phesphorus allowances should be approximately 1.5 times those for entrum. When the enforces are oblamed largely from egents computation of the total phesphorus may be masheding because phytin phosphorus may be poorly utilized untess the supply of vitamin D is adequate

Copper The requirement for copper for adults is about 1 to 2 mg daily Intants and children require approximately 0.05 mg for each kg of body weight. The requirement for copper is approximately one tenth that for iron

Joine The requirement for jodine is small probably about 0 002 to 0 004 mg daily for each kg of body weight, or a total of 0 16 to 0 30 mg y for the adult J specully unpertant in addressnes and pregnancy. The daily intake should approximate about 2 mg Pitamin Be Gro p Vitamin Be is essential for human nutrition daily for the adult

Polecin It were probable that cletary intakes of the order of less than a mg per day one be expected to cover any nutritional needs for folio need

3 total 18 needed by the human Partithene And The amount of pantothense and consumed daily with 2500 calones of good duet is approximately 10 mg Bieten Intestinal synthesis of this vitamin is extensive and tends to provent the development of deficiency oody and cannot be synthesized metabolically

daily oral dove of 1 mg. should be satisfactory
A suntable smount for the mother preceding delivery is 1 mg. administered parenterally this amount wil stabilize the prothroubin level of the infant until food is taken A suntable smount for the infant after birth is 1 ng for a single ilose In fact it has been shown that a single dose of 10 to 20 µg is adequate to cover the first 5 days. The daily requirement of the infant is approximately 1 µg lifamin A Available evidence warrants increased attention to the vitamin K incake of the mother during the latter part of pregnance

qurements less well known the allowance levels are considered to cover individual variations among normal persons in they live in the United States publected to ordinary environmental stresses common thereto. [bor a proper understanding of the foundations underlying this Tuble of Recommended In planning practical dictaires the recommended allowances can be attained with a variety of common foods also providing other nutrient re Jaily Monances reference should be made to the original source]

There calorie recommendations apply to the degree of activity for the reference man and woman described in the original text. I or the urban white coll it worker they are probably excessive In any case the calotie allowance must be adjusted to the netual needs of the individual as indirate | by his weight and height [Adjustment for cuvironmental temperature should also be made]

'The recommendations for infants pertain to nutrients derived primacily from cows milk or commercial milk preparations. There should be no question that human milk is the most describe source of nutrients for infants although expected intakes may not provule the recommended lovels of ertain nutrients og protein calcium and riboffavin microbiological procedures have been developed (particularly for the vitamins of the B group) based on the propagation of various bacteria, yeasts, or molds in special media in which one of these vitamins is the limiting factor

The conditions employed in the nonhological assay procedures, especially in the preliminary extraction or hydrolysis of the samples, are of course not necessarily equivalent to the conditions to which such foods may be subjected in the gastraintestinal tract. Therefore vitamin content as determined by chemical analysis may not always coincide with vitamin potency as measured by feeding tests Correlation of animal assays of foods with nonlinological assays suggests that the full content of vitamins is usually, but not always, biologically active under the experimental conditions employed The assessment of vitamins in foods in terms of their physiological availability for man is a field of investigation which needs more intensive exploration

It is important when using tables of the distribution of vitamins in foods, such as that presented in Appendix III, to bear in mind that most of these data are based on assays for total vitamin content rather than for available vitamin content in the physiological sense. For example, in tables spinach, whose "vitamin A" is due to the activity of carotenes, or butterfat, whose "vitamin A" is due to a mixture of carotenes and preformed vitamin A, are usually presumed to have been evaluated biologically, and whenever possible the interpretation of nonbiological

assays is made in the light of existing animal assay data

When referring to food-composition tables it must be recognized that uniformity is the exception rather than the rule Such factors as soil, chmate, season varietal differences and period of harvesting (which affect plant foods) or composition of the feed, age at slaughter, and duration of lactation (which affect animal foods), as well as storage, transportation processing "refining," and culinary losses, account for a several fold range of variation in the vitamin content of natural foods. Hence the values given in tables must be regarded as indicative of expected orders of magnitude rather than absolute vitamin contents

Under the provisions of the Food, Drug and Cosmetic Act, the Food and Drug Administration established standards of minimum daily require ment for certain of the major vitamins (and minerals) to serve as a basis for proper labeling of foods intended for special dictary uses. These values are presented as a guide to the interpretation of such labeling but are not to be confused with the table of recommended daily allowances prepared by the Food and Nutrition Board of the National Research Council reproduced in its latest (1953) revision on p 1108 The latter is intended to furnish "desirable goals toward which to strive in planning dicts and food supplies' It provides for variations in requirements due to sex, age, activity, and the physiological demands of pregnancy and lactation Individual needs may deviate significantly from these allovances owing to differences in body weight, glandular activity, gastrointestinal disturbances and many other conditioning factors

The term "major vitamins" is sometimes applied to thiamine, riboflavin macin ascorbic acid and vitamins A and D, for which fairly reliable knowledge exists as to human requirements, it is generally believed that if diets are devised to supply adequate amounts of these vitamins, sufficient quantities of all other vitamins (somewhat presumptuously ealled "minor") will be present. This view is questionable and merely reflects current ignorance as to the essentiality and quantitative requirements for such vitamins which, it is hoped, future research will correct

Whether a given food, as prepared for consumption, is a poor, fair, good, or excellent source of the vitamins or of other dictary essentials must be judged by the size of the usual portion and the frequency with which it is consumed. The Council on Foods of the American Medical Association* tentatively adopted the following standards for determining the significance of a source of a dictary essential:

 In general, when one-tenth of the day's requirement for an average man is furnished in a portion which can be easily eaten in one day, the food may be regarded as a "fair" source

2 When one-tenth of the day's requirement is contributed by an amount of the food which at the same time furnishes not more than 200 calories, the food may be classed as a "regod" source.

classed as a "good" source

3. When one-tenth of the day's requirement is furnished by a food which appears in the diet practically every day, and in which the portion contributing one-tenth of the essential furnishes not more than 100 calories, the food may be classed as "excellent"

4. When a food is not one which can be easily eaten in amounts to furnish one-tenth of the day's requirement, or is one eaten infrequently, or both, and the amount required for one-tenth of the day's allowance of the essential furnishes more than 200 calories, the food is a negligible or poor source.

VITAMIN A

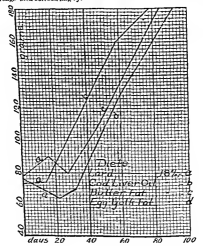
Physiological and Clinical Aspects of Vitamin A. Vitamin A was first recognized by the failure of rats to grow on diets lacking in natural fats and oils At about the time of cessation of growth the eyes become hemorrhagie, keratinized, and later infected This condition, known as serouhthalmia, is associated specifically with deficiency of vitamin A, hence the name antiophthalmic (or antixerophthalmic) vitamin (Fig. 271). Interruption of growth (Fig 272) and lowered resistance to bacterial infection are secondary results of a specific disturbance in the metabolism of epithelial membranes resulting from vitamin A deficiency. This is deduced from the fact that keratinization is observed in the sublingual and submaxillary glands, and in the respiratory, alimentary, and genitourinary tracts, as well as in the cornea and conjunctiva. Similar involvement of the oral and pharyngeal mueosa is responsible, at least in part, for the diminution of food consumption and consequent loss in weight. The formation of urinary and renal calculi observed after vitamin A depletion is probably also a sequel to the metaplastic changes in the epithelial membranes. Similar alterations in mucosal epithelium are followed by bacterial invasion and consequent infections of the sinuses, middle ear, and other areas. However, vitamin A has no specific or immunologic value in the treatment of the common cold. A form of dryness of the skin (xerosis) and a tollicular hyperkeratosis (phrynoderma) are among the

⁴ J. Am Med Assoc., 108, 1890 (1937).





Fig 271 Ophthalana is Two Stages or Strepfty (Left) Incrustation caused complete closing of eye (PigHt) Depulated and himorrhagic niea surroun hing eye

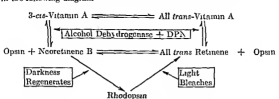


symptoms of avitaminosis A found in man as well as in animals. A variety of elin diseases characterized by these symptoms have responded to vitamin A therapy (see p. 1290). Suggestions have been made to name utamin A "anti-infective" or "antikeratine." but these terms have not been adonted because their implications are too broad Inantion accomnanying severe vitamin A denletini may contribute in no small degree to the lowered resistance to bacterial invasion. The primary specific effect of lack of vitamin A, however, is the "substitution of stratified keratinizing enithelium for normal epithelium in various parts of the respiratory tract, alimentary tract, eyes and paraocular glands, and the genito-urnary tract" (Wolbach)

There is evidence that lack of vitamin A may produce degenerative changes in the michin sheath Since the lack of vitamin A interferes with the process of ovulation, an adequate dictary supply of this vitamin is necessary for normal fertility. It has been demonstrated in rats that a liberal allowance of this vitamin is conducive to longevity

In addition to its role in maintaining the normal cuithelium of the eve and paraocular glands, vitamin A plays a critical part in the process of vision. The characteristic pigments of the retinal rods and cones are conjugated carotenoid proteins known respectively as rhodopsin and andonsin According to Walds they differ only in respect to their opsin or protein moieties (called sectopsin and photopsin, respectively) The specific carotenoid common to both is a cis isomer of retinene. The range and degree of sensitivity of the rods to light (scotopic vision) is determined by the absorption spectrum of rhodopsin, whereas that of the concs (photopic vision) is determined by the absorption spectrum of เดสิดทรเท

The chemical changes in the visual cycle involving rhodopsin are shown in the following diagram



In the pigment cpithelium of the rods a cis isomer of vitamin A alcohol is oxidized to the corresponding vitamin A aldehyde, retinene This reaction is reversible and is mediated by alcohol dehydrogenase in the presence of cozymase (DPN) The protein opsin, acting as an aldehyde trapping agent, combines stoichiometrically and spontaneously with retinene in the dark to form rhodonsin or visual number This is the basis of

Green and Mellanby Brit Med J 2 (91 (1998)

Nedder Porto Rico J Pr. Health and Trop Med 5 293 (1930)

Mald Harrey Lectures 41 117 (1945-7) Science 113 287 (1951)

dark adaptation Light induces a series of photochemical changes in rhodop in, beginning with bleaching of the purple pigment on the releas of the visual impulse, and ending with the formation of all-trans-retinen and its isomerization. For each molecule of retinene formed two sulfly dryl groups are liberated, and in the reverse reaction two free HS group are provided by open.

The resynthesis of rhodopsin is isomer-specific, requiring a cts isome of retinene (probably 3-cts or neoretinene B) for the formation of which the corresponding cts isomer of vitamin A is withdrawn from the circulation Equilibrium reactions involving the isomerization of trans-retinen or vitamin A to cts forms take place elsewhere in the body, probably i

the liver

The reversible synthesis of rhodopsin has been reproduced in ritro h
Hubbard and Wald's using the four-component system vitamin A (nature

or as isomers), cozymase, alcohol dehydrogenase, and open Since some vitamin A is lost in the visual excleded deficiency of the vitamin results in a diminution of visual actuity, i.e., the ability to see in subdued light. This condition is known as nyctalopia or might himdness and is the earliest sign of vitamin A deficiency in man, though it may result from other causes. Dark adaptation is the term applied to the adjustment of the usual threshold to darkness after exposure of the exist to light. Quantitative measurement of the visual threshold—ie, the light intensity required to clicit a visual sensation—is made hy photometric instruments which measure the rate of restoration of visual actury following stimulation by a light source of standard intensity.

Severe outbreaks of vitamin A deficiency in humans are rare and of limited geographic distribution. The incidence of ophthalmia is relatively high for example in China, Lahrador, and India. An acute epidemic among Danish children during World War I was attributed to the substitution of margarine for butter, which was largely exported. The fortification of margarine with vitamin A is now a common practice in most countries where it is sold.

Twenty to 25 units of vitamin A per kg body weight suffice to maintain a normal visual threshold in most species studied. This amount will also maintain the normal estries cycle in the rat, although storage in the liver does not occur until approximately four times this level is reached. Liheral allowances of vitamin A, among other factors, tend toward optimum, as distinguished from adequate, nutrition (see p. 1074). In experiments on the rat Sherman and his associates have shown that by feeding several times the dose of vitamin A sufficient to prevent deficiency symptoms, the reproductive period of females as well as the longevity of both sexes could be prolonged.

The efficiency of absorption of vitamin A in different species is influenced by the nature and quantity of the diluent and by the state of the

VITAMEN A AND D CONTENT OF OILS FROM FISHERY SOURCES HAVING COMMERCIAL IMPORTANCE IN THE UNITED STATES AND ALASKY.

F		S	surce of C	a	Vstamin A USP Units per g of Oil		Litamii	
Common Name	Scientific Name	Area	Organs	Per Cent of Body Il eight	Oil an Organs Per Cent	Range	Aver	D IU per g of Oil
Soupfin shark	Galeorhinus zyop-	Pacific	Liver	10	55-68	45 000- 200 000	120 000	5-25
Coupfin shark	Galeorhinus zyap-	Pacific	Liver	10	65-72	15 000- 40 000	32 000	5-25
Gravish (dog fish)	Squalus suckleys	Pacific- Alaska	laver	10	67-72	2 000- 20 000	3 000	5 25
Gravfish (dog fish)	Squalus suckleys	Pacific US	Liver	10	50-70	8 000- 25 000	14 000	
Halibut	Mappoglossus happoglossus	Pacific	Liver	1-3	8-27	20 000- 160 000		1000-000
Hall ut	Happoglossus happoglossus	Pacific	Vis ecta	^ 5–5	2-5	70 000- 700 000	200 000	
af lefish	Anoplopoma fim brsa	Pacific	Luet	2 2 5	10-26	50 000- 190 000	80 000	
Fablefish	Anoplopoma fim brea	Pacific	V15- cera	3-4	5-12	90 000- 250 000	1º5 000	
L ngcod	Ophiodon elong- alus	Pacific	Liver	115	1	40 000- 550 000	l	1000-6000
Langeod	Ophioden clong	Pacific	V1S ceta	18-3	4-15	10 000- 175 000		100-200
llamn erhead shark	Sphyrna tygarna	Pacific Atlantic			30-40	30 000- 120 000	50 000	
Albacare tuns		Pacific	Liver	1 5-2	7-0	\$0 000 10 000	25 000	250 000
Bluefin tuna Bonito	Thunnus thynnus Sarda chilensus	Pacific Pacific	Liver		4-6	25 000- 100 000	75 000	70 000
Swardfish	Yuphias aladius	Pacific	Laver	1 4-2 6	4-12 8-35	15 000- 50 000-	35 000 35 000	50 000
Cod	Gadus callarius	Atlant c		3-5	20-60	400 000 1000-6000	2000	25 000 100-600
Hal but	Happoglossus happoglossus	Atlantic	Liver	1 5-2 5		1100-6000		1000-5000
Herring Menhaden	Clupea pallasis Brevoorlia tyran	Pacific Atlantic	Body Bods		5 °0	50-300	90 500	25-160
	,							

^{*} Butler Commercial Fisheries Rev April 1946 p 13

vitamin, 1e, whether carotene, preformed, free, or esterified ^{11a} The recommended daily allowances of vitamin A for various sex and age groups are incorporated in the table on p 1108 These values are based on a consensus obtained under the sponsorship of the National Research Council An outline of the climical symptoms of vitamin A deficiency is given in the American Medical Association svillabins on p 1290

Storage of Vitamin A. When young animals, previously fed a diet contuning vitamin A, are deprived of this vitamin, they continue to grow

¹¹a Week and Seviene J Nutration 39, 233 251 (1949) 40 563 (1950)

for a period dependent upon the amount of vitamin A which they have stored. The storage amounts is a the age of most ripid growth. The vitamin is relatively gir iter in young animals, i.e. at the age of most ripid growth. The vitamin is stored principally in ester form in the liver and kidneys and to some extent in the lings. At high levels of intake rats store more vitamin A in the liver than in the kidneys, but at low intake levels the reverse is true. I we storage is the basis of a bioassay for vitamin A.

Distribution of Vitamin A 1 Preformed vitamin A occurs only in lipides of animal origin whereas carotenoid precursors occur in the vegetal le knigdom where they are synthesized. The fat of milk may contain both carotenoids and vitamin \ in variable proportions depending on the animal s ration and the extent of conversion of the former into the latter The chief natural foods of vitamin \ value are butterfat (hence milk eream cheese etc.) egg yolk hver and pigmented and leafy vegetables (e g carrots tomatoes pimientos apinach lettirec alfalfa) Vegetable oils as a class are deficient in vitamin \ although red palm oil may contain as much as 0.2 per cent of carotenoids. The hier oils of certain species of fish including the shirk (notably the soupfin species) swordfish halibut mackerel ete are extremely high in vitamin \ some oils I cing several hundred times as rich as codiner oil one of the earnest known and most abundant sources of both vitamins V and D. There are great variations within species due to storage with age sex and reproductive phase marine nutrition temperature ete Natural fish hier oils are refined or concentrated and blended to standard potencies for thera neutre or food use

Certain unicellular marine organisms are able to synthesize vitamin A. These provide food for marine plankton, which are consumed by caplin and other small fish, which in turn are consumed by the larger species. The vitamin is also present to a smaller extent in the visceral and body oils of the cod, salmon, and certain other fish.

In the plant world vitamin \ activity being due to earotenoids 15 closely associated with pigmentation the green outer leaves of lettuce are richer than the white center leaves green tips of asparagus are richer than bleached tips sweet potatoes are richer than white pota toes yellow corn than white palm oil (red) than coconut oil The milk of cows on green pasturage is higher in vitamin A than is the milk of stall fed animals to direct quantitative relationship exists between the color of butter or of fish liver oils and vitamin A potency since in animal fats such activity is due principally to the colorless preformed vitamin (Com mercial butter is often artificially colored) The yellow pigment carotene possesses the physiological activity of vitamin A by virtue of its con version into the vitamin in the intestinal wall. Hence carotene or rather certain carotenoids are provitamins or precursors of vitamin A In ad dition to the carotene isomers (see below) other earotenoids which possess, vitamin A activity in animals are eryptoxanthin of vellow corn and paprika myxoxanthin and apl anin of algae

The synthes s of provitamins A in plants has been shown to be acceler ated by hight radiations especially of short wavelength although the

² For vitam n A values of foods see Append z III

ultimate amount of vitamin formed is not increased. There is no evidence that the animal body can produce vitamin A de novo, except in the case of the pigeon. Although synthesis of carotenes seems to parallel chlorophyll formation, the latter pigment does not possess vitamin activity.

The rate of intestinal absorption of vitamin A and its precursors varies in different species of animals and is influenced by the nature and content of fat in the duet Excessive quantities of mineral oil inhibit absorption. The efficiency of converting carotenoids into vitamin A likewise varies with the particular compound as well as with the species; in general, rodents are most efficient, pigs and cattle less, and cats least, while the relative capacity of the buman infant or adult in this respect remains to be determined

Chemistry of Vitamin A. β -Carotene has been shown by Kuhn and Karrer to have the following structural formula:

In the animal organism cleavage occurs at the central double bond with the formation of the alcohol, vitamin A

¹⁵ Kon and Drummond Biochem J., 21, 632 (1927).

Hunter and Williams¹⁴ claim to have effected this conversion by oxidation with hydrogen perovide to give vitamin A aldehyde, and subsequent reduction to the alcohol. In the liver and other storage depots of the animal organism vitamin A exists 11 the form of fatty acid esters. The palmitate and acetate are produced commercially from either natural or synthetic sources.

It will be noted that the β -carotene molecule is symmetrical, containing two β -ionone rings, designated β^1 and β^2 respectively, bridged by a chain of four isoprene (CH₂=C+CH=CH₂) units. The α and γ isomers differ

cilia in respect to the configuration of only one of the rings, the molecular formula of all of these isomers being $C_{40}H_{46}$. In place of the rings indicated above for β -carotene, certain other carotenoids differ only in respect to the rings as shown in the following tabulation.

At least one intact β -ionone ring is essential for vitamin A activity, since isomers or derivatives in which the only change is in both rings (e.g.,

¹⁴ Hunter and Williams, J. Chem Soc., 554 (1945), Hunter Nature, 158, 257 (1946)

lycopene or vanthophyll) are mactive. The conjugated double bonds in the side chain (or in the carotenoid bridge) make possible a large number of cis trans isomers ¹⁸ However not all of these stereoisomers exist in nature or have been produced synthetically, owing to steric hindrances. Twenty cis trans isomers of β carotene are possible and even larger numbers in the case of the asymmetrical compounds like α - or γ -carotene. The stereoisomers are differentiated by their spectral absorption characteristics in both the visible and ultraviolet wavelengths and by their chromytographic adsorption affinities.

It has been postulated that because of steric interference, only two of the four double honds in the side chain of vitamin A assume cis trans configuration, yielding four stereoisomeric forms viz all-trans, 3-cis-, 5-cis-, and 3,5-di-cis-vitamin A. The isomers found in liver oils are principally all trans (Fig. 273) associated with a cis isomer (probably 5-cis) which has been crystallized and designated "neovitamin A" (Fig. 274). Despite the functional differences between these isomers in the visual cycle (see p. 1113) the growth-promoting activity of neovitamin A for the rat appears to be approximately 80 per cent of that of crystaline all trans-vitamin A."

The approximate vitamin A activity for rats of certain important natural carotenoids relative to β carotene is as follows

β-Carotene (all-trans)	100
a Carotene (all Irans)	53
y Carotene (all trans)	27
Cryptoxanthin (all trans)	57
Neo-β earotene U(mono-cis)	38
Neo-8 earotene B (di-cis)	53
Lycopene	Ö
Nanthophyll	0

β Carotene is approximately twice as potent biologically as its naturally occurring isomers. It was believed that fission of the carotenoid molecules occurs at the central double bond yielding two molecules of vitamin A there is evidence, however, that asymmetrical breakdown to the active β ionone fragment of the molecule can also occur.

Since intestinal absorption, stability, and efficiency of conversion affect the biological activity of carotenoids the estimation of vitamin A potency of plant foods from the relative content and activity of the individual pig

ments is not always reliable

The previous international standard of vitamin A activity, β caretene has been replaced by crystalline vitamin A acetate. The International or USP unit is now defined as 0.30 $\mu_{\rm B}$ of vitamin A alcobol equivalent to 0.344 $\mu_{\rm B}$ of the acetate. For the estimation of carotene the former stand ard has been retained, 0.6 $\mu_{\rm B}$ of pure β -carotene is considered to have the

¹⁵ For reviews of stereoisomerism of the carotenoids and vitamin A see Zechmeister 1 tiamins and Hormones 7-57 (1949). Karrer and Junker. Carotenoids. New York. Pleevier Press Inc. 1950.

Robeson and Baxter \aiure 155 300 (1945)
 Hartis Ames and Brinkman J Am Chem. Soc 73 125° (1951)



FIG 273 CRYSTALLINE VITAMIN A ALCOHOL (HIGH MELTING)
Courtes Distillation Products Inc. Rockester, N. Y.



FIG 274 CRYSTALLINE NEOVITAMIN A. Courtesy Distillation Products Inc. Rochester N Y

activity of one unit of vitamin A. It has been found however that the "erude carotene" of many plant foods (i.e., the pigment in the petroleum other extract which absorbs light at $420~\mathrm{m}\mu$) has a vitamin A activity of

about 1 USP unit per ug. Vitamin A alcohol forms pale vellow ervstals melting at 63-64° C. when deposited from ethyl formate solution or at 7-8° C. when methyl alcohol is the solvent. It forms esters but is not precipitated by digitonin. providing a means of separation from cholesterol which likewise occurs in the nonsaponifiable fraction of animal fats. The vitamin is distillable in racuo, thus forming the basis for a process of concentration from fish liver oils: viz. Hickman's method of molecular distillation. Vitamin A forms colored compounds with certain condensing agents, one of which. antimony tricbloride, produces an evanescent blue color (Carr-Price reaction) of sufficient intensity to afford a measure of its concentration. This color absorbs light maximally at 620 mg, in contrast with the more stable greenish-blue color produced with earotene, which shows an absorption maximum at 590 mu. For the analysis of foods or animal tissue containing both factors, the ShCls reaction necessitates either correction for the differences in rate of color development and fading, or preferably, pre-

liminary separation of the carotenes from preformed vitamin A.
Vitamin A is characterized by an absorption band in the ultraviolet
with a maximum at 328 ma. This forms the basis of the spectrophoto-

metric method of estimating (preformed) vitamin A content.

In the absence of air, vitamin A is quite stable at high temperatures, but when exposed to air or ovygen it is readily destroyed Aeration of codiver on the 12 hours at 100° C. ovidizes the vitamin A, leaving the antirachitic potency unimpaired. Vitamin A appears to be more resistant to ovidation in its natural environment, probably because of the presence of protective antiovidants such as tocopherois, prospholpides, and possibly other agents. Certain phenohe substances, like hydrogunone and pyrogallol, are even more effective antiovidants, but their use is not permitted on toxicological grainds. Light, especially in the ultraviolet range, exerts a destructive influence upon vitamin A; hence the practice of dispensing lish liver oils in dark bottles and of conducting analytical operations in amber or red glassware.

DETERMINATION OF VITAMIN A

In estimating vitamin A in natural products a distinction must be drawn between vitamin content and vitamin potency. Vitamin A activity is derived principally from the physiologically available amounts of preformed vitamin A, which may be present as free alcohol or as esters, and of the carotenoids, which vary in their individual potencies. Furthermore activity is influenced by the nature and content of fat in the diet and by the physical state of the carrier; for example, fine dispersions of vitamin A in aqueous media are absorbed more rapidly than is the vitamin in oily media.

Because of this distinction between vitamin A content, in the chemical sense, and biological activity of true vitamin A and related compounds, the bioassay continues to play an important function, especially in the evaluation of foods whose vitamin activity is of multiple origin. In products whose vitamin Λ activity is derived from less complex sources, such as the crystalline alcohol or its esters, high-potency fish liver oils of β -carotene preparations, it is only necessary to establish correlation between values for vitamin content obtained by physical or chemical methods and the bioassay values to justify the use of the nonbiological procedures Because of the small size of sample available, the analysis of blood and tissues for vitamin Λ is limited to the nonbiological methods

CHEMICAL METHODS

Colorimetric Methods for Preformed Vitamira A and Provitamina A. The caroteness are determined colorimetrically preferably after isolation by saponification, extraction, adsorption, and elution Prediction of biological potency from the content of carotenoids is complicated by their individual differences in activity and absorbability. Chromatographic separation of the carotenoids and their identification and estimation from spectrophotometric absorption data afford a more accurate but involved basis for estimating the vitamin A activity of plant carotenoids.

Various color reactions have been proposed for the estimation of preformed vitamia A, especially in liver oils 18 The most widely used reaction is that of Carr and Price 18 When a solution of antimony trichloride in chloroform is added to a dilute solution of a vitamin A-bearing oil, a blue color appears which soon reaches a maximum intensity and their rapidly fades or changes to reddish-brown or other colors, varying with the individual oils Under carefully controlled conditions the blue color persists long enough to make accurate readings possible Glycerol 1,3-dichlorohydin activated by distillation over antimony trichloride is also used as a colorimetric reagent for vitamio A, greater color stability being claimed for the reaction product 29

Antimony Trichloride (Carr-Price) Reaction: Into a dry test tube introduce 0.2 ml. of a 20 per cent solution of codliver oil in chloroform. Add rapidly 2 ml. of a saturated solution of antimony trichloride in chloroform. Observe the color changes.

Determination of Carotene (Provitamin A). β-Carotene occurs in certain plant sources (e.g., carrots, alfalfa) relatively free from other carotenoids such as cryptoxantin or lycopene in these cases it may be determined by methods involving selective solvent extraction (phasse separation) Mixtures of hexane and acctories, methanol, or diacetone alcohol are widely used, since they permit partition of the carotenes from xanthophyll, chlorophyll, or other mactive pigments. When present

with other carotenoids, β-carotene is preferably separated by chromatography of a

Extraction. In the case of animal or dairy products containing fat, saponification and extraction is necessary. For this purpose the isolation of the unsaponifiable extract as described for vitamin A on p.1124 may be employed, but the residue after evaporating the ethyl ether should be taken up in an appropriate volume of hexane. Drued plant products or feed mixtures may simply be extracted for one hour under reflux with a mixture of acetone and hexane (3 + 7), using about 30 ml. per 2 g. of sample. After cooling, the extract is filtered through anhydrous sodium suffate into a 100-ml. volumetric flash, and made up to volume with hexane rinsings.

Other extraction methods have been described for special cases, e.g., those requiring cold extraction to avoid isomerization.

Chromotography. In view of the predominance of \(\beta\)-carotene in most animal tissues or foods intended for human use, the method to be described, which actually measures total or "crude" carotenes, will generally be found satisfactory. For procedures effecting more complete resolution of the carotenes and their isomers the reader is referred to the literature."

Insert a chromatographic adsorption tube (22 × 175 mm.) into a suction flask and plug the outlet orifice with a bit of glass wool. Introduce a mixture of equal parts of activated magnesian and Hyflo Supercell to a depth of 15 cm. with the aid of suction and a flat-surfaced tamping device; gently press the adsorbent down to a depth of 10 cm. Cover with a 1-cm. layer of anhydrous sodium sulfate.

Under continuous suction pour the extract into the adsorption column. (Always keep the top of the column covered with a layer of solvent.) Follow through with 50 ml. of actone-hexane (1 + 9) to develop the chromatogram. Collect the cluste, which contains all the carotene, leaving the xanthophylls, chlorophyll, and oxidation products in the column. Transfer the partially evaporated cluste to a 100 ml, volumetric flash and make up to volume with acctone-hexane.

Spectrophotometry. The solution should have an absorbancy corresponding to 1-4 \(\rho_8\), of carotene per mil. Read the optical density in a spectrophotometer at 436 mm. Convert to "total carotenes" by means of a calibration chart prepared from graded concentrations of a standard solution of crystalline \(\rho_8\)-carotene="i" or calculate from the following equation:

$$\frac{(-\log T) \times v \times 100}{196 \times L \times C} = \text{mg carotene per } 100 \text{ g sample}$$

where $(-\log T)$ is the optical density, v the final volume of cluate (100 ml), L the cell depth in cm, and C the weight of the original sample

¹¹ Beadle and Zscheile J Biol Chem., 144, 21 (1942). Zechmeister and Polgar J Am Chem. Soc., 64, 1856 (1942)

²¹ Bickoff, et al. J. Assoc Official Agr. Chemists. 31, 633 (1948). 32, 766 (1949). These authors employ hydrated lime as an assorbent to separate the carotene isomers.

Micron Brand No 2642 Westvaco Chlorine Products Co , Newark, Col

^{*}Johns-Manville Co, Cheego III
*The International Standard obtamable from USP Reference Standards, 46 Park Ave, N Y 16 Pure β-carotene or a maxture (90 per cent β, 10 per cent α) as supplied by General Boochemicals Inc. Chargin Falls Ohos may be used β-carotene may be purified as follows Dissolve 100 mg β-carotene in 2 ml chloroform, and reprecipitate with 20 ml methanol Filter, wash with a few drops of methanol, and dry an a vacuum desected.

Determination of Preformed Vitamin A (Colorimetric Method of Out., Melnick, and Pader): Principle. The I has color produced when a term A reacts with antimony treliborate in chiroform solution is inexaired photometrically As internal standard (increment) is included to compensate for the effect of inhibitors of accelerators on the color development.

Procedure: Preparation of the Unsaponlfiable Fraction. Perform all operations in amber glassware. Into an Frienmeyer flask transfer a weighed sample, containing preferably at least 50 USP units of vitamin A. The sample mar he weighed by difference from a weighing bottle furnished with a rod and stopper. Saponify by refluxing on a holling water hath for 0 5 hr. with freshly prepared 0 5 N alcoholic potassium hydroxide, using 15 ml. for each gram of sample solids. Cool, and transfer to a separatory funnel, adding an equal volume of water as a wash. Fatract four times with 75 ml, of freshly redistilled ether, and discard the aqueous phase. Wash the combined ether extracti once with 50 ml, of water, once with 25 ml, of 0 5 N aqueous potassium by droxide, and then with 50-ml, portions of water till the last washing gives no color with phenolphthalein, Dry the ether extract with anhydrous sodium sullate and evaporate to dryness on a water bath, removing the last few mililiters at room temperature with a stream of nitrogen. Dissolve the residue immediately in sufficient purified" chloroform to produce a concentration of from 5 to 15 USP units of vitamin A per ml. A turbid chloreform solution may be clarified with anhydrous sodium aulfate.

Colorimetric Procedure. A direct-reading photoelectric colorimeter with a 620-m; filter is used for the measurements. Set the instrument at 100 per cent transmittance with a solution containing 1 ml. of chloroform and 9 ml of antimony trichloride reagent. To another tube add 1 ml of the test solution and 9 ml. of chloroform, and read (A) To a tube containing 1 ml of chloroform extract of the sample, add 9 ml. of antimony trichloride reagent rapidly from a blow-out pipet and measure the maximal blue color as indicated by the full swing of the galvanometer usually attained within 4 seconds (B). To another tube containing a 1 ml. aliquot of the test solution, with 2 mlcropipet add 0 1 ml of 2 vitamin A standard* containing 10 USP units (of 3 µg.) of vitamin A followed by 9 ml of the antimony trichloride reagent Measure the maximal color (G)

CALCULATIONS Convert galvanometer readings, G_t expressed in per cent transmittance, to photometric densities, PD_t as follows

$$PD = 2 - \log G$$

Then,

$$\frac{B-A}{101(-B)} \times 10$$
 USP units (or 3 µg) × dilution factor

⁼ LSP units (or µg) of vitamin A per g of sample

[&]quot;Oser Melnick and Pader Ind Eng Chem Anal Ed 15 724 (1943), Oser Melnick. Pader Roth and Oser 2nd 17 559 (1945)

^{**} Wash reagent grade chloroform three times with an equal volume of water and dro over anhydrous sodium sulfate Redistil on the day used

TA 20 per cent solution of autumony trachlorade in dry chloroform Filter if solution is turbed

The USP Reference Standard Solution of crystalline vitamin A acetate may be employed as the standard the results are expressed in terms of vitamin A sloohol

Spectrophotometric Method.³¹ Since vitamin A is characterized by selective absorption in the ultraviolet, the intensity of absorption at the maximum (Fig. 275) serves as a measure of vitamin A content. Inasmuch as this region of the spectrum is beyond the visible range, spectrophotometric equipment is necessary. Adaptations of this method have been particularly useful in the assay of fish liver oils where the vitamin A

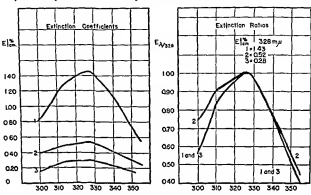


FIG. 275 ADVANTAGE OF PLOTTING EXTINCTION RATIOS RATHER THAN EX-TINCTION COEFFICIENTS IN PRESENTING ULTRAVIOLET-ABSORPTION CURVES OF VITAMIN A SQLIVINGS.

1. Vitamin A acctate in ethyl laurate 2 Oxidation of (1) by air. 3 Dilution of (1) 1.5 with ethyl laurate

activity is due entirely to the vitamin per se. Greater accuracy is obtainable by the removal of interfering substances present in the saponifiable fraction, particularly in oils containing less than 15,000 units of vitamin A per gram.

The method involves the comparison of the absorption of ultraviolet radiation through the test solution containing vitamin A, with that absorbed by the solvent alone. Results are usually expressed in terms of the extinction coefficient! derived as follows:

$$\Gamma_{1\text{em}}^{1\text{e}} = \frac{1}{cd} \log_{10} \frac{I}{T}$$

in which $E_{lm}^{1/2}$ is the extinction coefficient, c is the concentration (in per cent), d the thickness of the cell (in cm.), and I and T the intensity of the

¹¹ For treatments of this subject, see Chapter 23, also Morton The Application of Absorption Spectra to the Study of Viamous, Hormones and Coenzymes 2d ed. London, Higer, 1942, Chapter 4 in Wokes Applied Biochemistry, Baltimore, Win Wood and Co (The Williams & Wilkins Co), 1937, and U.S Pharmacopena XIV, 1950, pp 756-8
2" "Specific absorbane," in the terminology of modern optical physics

"incidental" and transmitted light, respectively. For a discussion of the derivation of this equation see p. 512 et seq.

To be valid as a basis for estimating vitamin A content, the absorption curve from which the extinction coefficient at the peak (325 mµ) is taken should he characteristic of the pure vitamin A absorption curve. Irrelevant absorption due to exidation products of the vitamin or to other inactive substances may be evidenced by distortion of the vitamin a curve. This is more readily recognized when the curve is plotted without regard to concentration of the vitamin, i.e., on the assumption that the extinction coefficient at the maximum is equal to unity. The value at any other wavelength thus becomes an extinction ratio, i.e., the ratio of the extinction coefficient at that wavelength to that at the maximum (Fig. 275)

The conversion of the extinction coefficient (or "E value," as it has been abhreviated) to vitamin A units is effected by multiplying by a conversion factor derived from observed ratios of biological potency to E value in fish liver oils. In practice such conversion factors have been found to vary over a wide range in different laboratories, owing to differences in the oils themselves, as well as to the errors inherent in both the biological and instrumental methods employed. On the bass of improved analytical techniques as well as theoretical considerations, it is now commercial practice to regard the factor 1900 multiplied by the extinction coefficient at the maximum for the typical vitamin A curve as giving the best estimate of vitamin A unitage.

Determination of Preformed Vitamin A (Spectrophotometric Method): Principle. Vitamin A may be determined by measurement of the extinction coefficient of the characteristic absorption band in the ultraviolet region of the spectrum, whose maximum is in the region 323-328 mg.

SPECTROPHOTOMETRIC DETERMINATION AND CALCULATION Using quartz curettes. read the absorbance of the isonronand solution at 310 325 and 334 mg. The median was along the absorption maximum of pure vitamin A alcohol in isopropanol whereas the higher and lower are the wavelengths at which the absorbance is \$- (i.e. 0.857) of the maximum. The absorbance rates Increase and Assorbe are frequently found to deviate by more than 1 per cent from 0.857 due to the presence of non attamin A-absorbing materials. Assuming that such irrelevant absorption is linear within parrow wavelength limits (e.g. % above and below the peak) the correction procedure devised by Morton and Stubbs may be applied 34 In simplified form 35 this has been adonted into official USP and AO C vitamin A assays

A (corrected) = 7.4 m - 2.625.4 m - 4.375.4 m

in which A is the absorbance at the indicated was elength

1(corrected) \times 5.7 = 42 vitamin A (per g or other sample unit) in which L is the length of the absorption cell in em. and C the fraction of the sample unit per ml of ealution read

us Vitamin A × 3.33 = International (USP) units

Comment This method is applicable chiefly to fish liver oils where the vitamin is present in comparatively high concentrations. Though a more accurate value is obtained by conducting the determination on the unsaponifiable fraction, oils of high potency may often be assayed with httle error by dissolving them directly in isopropanol. For the analysis of foods that have been enriched with vitamin A. a blank test of an un fortified sample or of a sample in which the vitamin A has been destroyed by selective ultraviolet irradiation36 is advised

Agreement between the basically unrelated spectrophotometric and colorimetric procedures may be regarded as highly presumptive of the accuracy of the nonbiological determination. In fact the official assays for vitamin A-bearing materials in the US Pharmacopeia and the Association of Official Agricultural Chemists specify that for the spectrophotometric assay to be valid the colorimetric test must agree within a ratio of 1.00 to 1.30

Determination of Vitamin A and Carotene in Blood Serum (Calorimetric Methad of Dann and Evelyn) " Principle Vitamin A and carotene are extracted from serum with ethyl ether after saponification with alcoholic potassium hydroxide Carotene is determined colorimetrically in the extract by photometric measurement of the light absorption of a choloroform solution at 440 mm and preformed vitamin A by measurement of the light absorption at 620 ms of the blue color produced by reaction of the vitamin with antimony trichloride

⁴⁴ Morton and Stubbs Biochem J 41 525 (1947) 42 195 (1948)

³⁵ Oser Anal Chem 21 529 (1949)

[&]quot;Lattle Ind Eng Chem Anal Ed 16 288 (1944)

"Dann and Evelyn Brochem J 32 1008 (1938) Vitamin V may also be determined in plasma or serum by the following procedure which avoids saponification. Add to the plasma slowly while slaking an equal volume of 95 per cent ethanol followed by 2 volumes of petroleum etler (b. pt. 40-60° C.) Stopper tightly and shake for 10 minutes. Centriluge one minute at low speed. Evaporate an adaquot of the petroleum etler layer on a water. bath (10° C) in a stream of N2 leating finally at 70° C for a few seconds Tle residue is taken up in etl anol free chloroform for colorimetric determination (himble J Lab Clin Wed 24 1055 (1939) Yudkin Biochem J 35 551 (1941))

Procedure. To 10 ml. of serum in a small gask add an equal volume of 95 per cent ethyl alcohol and 2 ml. of a 60 per cent aqueous solution of potassium hydroxide. Boil for 3 minutes. Pour the mixture into 10 ml. of water in a separatory funnel, and wash the flask with two 15-ml. portions of water, followed by two 25-ml. portions of ether, adding the washings to the separatory funnel. Shake the funnel vigorously for I minute, then allow the phase to separate. Discard the aqueous (lower) layer. Wash the ether phase by shaking vigorously with 10 ml. of water, then twice gently with 25 ml. o water. Filter the ether solution through a layer of anhydrous sodium sul fate on a sintered-glass filter. Wash the residue with 20 ml. of ether, com brings the wash with the filtrate. With the ald of a stream of nitrogeo evaporate the ether solution to dryness on a hot water bath. Take up the residue in 10 ml. of chloroform in the absorption cell of a photoelectricolorimeter.

Vieasure the light absorption of the solution at 440 m μ , setting the instrument at 100 per cent transmittance with pure chloroform. Calibrate the instrument by measuring the absorption of standard solutions of C.P. β -carotene in chloroform containing 0.2, 0.4, 0.6, 0.9, 1.2, 1.5, 1.8, 2.1, 2.5, and 3.0 μ S, per ml. Plot photometric densities against concentrations of carotene. From the graph, determine the concentration of carotenolds per ml. in the chloroform extract of the unknown, and multiply this value by 100 to obtain the concentration per 100 ml. of serum.

Place the cell in a hot water bath and evaporate off the chloroform with the aid of a stream of nitrogen. The latter provides an inert atmosphere and prevents oxidation of vitamin A. Take up the residue in 1.7 ml. of chloroform. Place the cell in the photoelectric colorimeter equipped with a 620 mm filter, set at 100 per cent transmittance with 0 ml. of a solution containing 1.7 ml. of chloroform and 8.3 ml. of antimony trichloride reagent (see p. 1124). Rapidly add 8.3 ml. of antimony trichloride reagent (see p. 1124) and determine the maximal extinction. Calibrate the instrument by conduction the test on pure solutions of vitamin A in chloroform containing 0.5, 10, 15, 20, 30, 40, 60, 80, and 100 sg. per 1.7 ml.

Calculation Plot photometric density against cooccutation of vitamio A per 17 ml. Determine the concentration in the chloroform extract of the unknown, and multiply this value by 10 to obtain the concentration of apparent vitamin A per 100 ml of serum. This value must be corrected for the interference of carotenoids which react with antimony trichloride to produce a blue pigment. The color produced by 1 μ g of preformed vitamin A is approximately equal to that of 20 μ g of β -carotene Determine the true concentration of vitamin A by means of the formula

$$A = \frac{\ell^*}{20} = \mu g$$
 vitamin A per 100 ml of serum

where A is the concentration of apparent stamm A and C that of carotene, both expressed per 100 mi of serum

Interpretation. The normal vitamin A level for blood serum is 15 to 60 pg (approximately 50 to 200 LSF units) per 160 ml 1 ng grincipal carotionois in blood serum are carotione and xanthophyll, which are present in a relatively fixed proportion. In Linder these conditions, measurement of

total carotenoids, which is necessary for the determination of total vitamin A activity, would be a good practical index of carotene content Chromatographic analysis, however, has demonstrated that the true carotene content varies from 10 to 50 per cent of the total carotenoid 40 The normal total carotenoid range in blood serum is 100 to 300 μg expressed as β carotene

BIOLOGICAL METHODS

Since no single chemical or physical method for estimating vitamin A activity is sufficiently specific for all compounds and mixtures possessing vitamin A activity, the biological assay remains fundamentally the most reliable means for determining vitamin A potency and is the basis against which the more precise nonbiological procedures must be explusted. Many of the limitations of bioassays are removed when they are planned to measure not only the response of animals to dosage of the test material but to known dosage of some standard of reference. By halancing the distribution of test animals among assay and control groups with respect to strain, litter origin, sex, age, and hody weight, and standardizing tho environmental factors, diet, feeding, dosing conditions, etc. it is possible to reduce the error of biological assay to a minimum. The assay can be set up according to a statistical design to permit control of these variables. and also to provide a measure of the limits of precision of the estimate and the extent to which individual factors such as sex, litter, etc contribute to the total variance An essential feature of hioassays in which responses are quantitatively graded to dosage is that the slope of the doso response curve (plotted logarithmically) must be reasonably parallel to that of the reference curve Otherwise the standard and unknown prepa rations are likely to be qualitatively different, e.g., the unknown may be contaminated with a substance that is touc or inhibits absorption. The effect of departure from parallelism may be measured in an assay designed to permit factorial analysis of variance

The curative metbod for assaying vitamin A is based on depleting young rats of their reserves of this vitamin and determining the relative growth responses of groups of these animals receiving daily supplements of the test material or of the vitamin A standard The basal due fed throughout the depletion and assay periods should be complete with respect to all nutrients required by the rat evcept vitamin A Serious error may be caused by failure to include in the basal vitamin-A-free duet a sufficient amount of vitamin E (tocopberol), since the responses of the animals may be influenced by differences in the antioudant content of

the assay and reference materials

41 XIV th Revision

The US Pharmacopera¹¹ method for vitamin A is a "rat-curative" assay and is most generally employed for the evaluation of food and pharmaceutical products As described, bowever, it is a minimal-potency assay (i.e., it provides for one assay and one reference group), and therefore permits only the statement that the assay material contains "less

[&]quot;Wohl Dictotherapy Clinical Application of Modern Autrition Philadelphia W D Saunders Co. 1945 p. 228

than" or "not less than" the unitage of vitamin corresponding to the administered dosage level. The method may be adapted to quantitative estimation of actual potency by feeding two or more dosage levels of the assay material and a similar number of dosage levels of the standard of reference (USP Reference Vitamin A Standard). The dosage interval—i.e., the ratio of each dose to the next lower dose—is constant in both assay and reference groups. The resulting data may he computed according to the analysis of variance, as illustrated by the example in Appendix VI. 40 give an estimate of potency, the standard error of this estimate, and, most important of all, an objective indication of whether the assay is a valid hasis for such estimation.

Biological methods have been described for vitamin A based on prevention rather than cure of the deficiency in rats, on keratinization of vaginal epithelium (which occurs at about the twenty-third day of depletion in young rats), and on the colorimetric determination of the vitamin A stored in the livers after controlled feeding periods. The latter procedure is less adaptable than the curative rat-growth method to quantitative study since it requires the administration of massive doses of the vitamin

The U.S. Pharmacopela hiological assays for vitamins A and D are given in full on pp. 1261 and 1264, respectively.

THE VITAMIN B COMPLEX

Since the vitamin concept was first postulated, the initial group of three vitamins has expanded to more than 20 hy virtue of the discovery of the multiple nature of previously known vitamins or the recognition, from hiological experiments, that as yet unknown nutrients must evist. Most outstanding in this regard has been the hranching out of the vitamin B complex, a group of water-soluble factors usually associated in nature with thiamine (the original vitamin B), rich sources heing liver, yeasts, and hrans The possibility that the "antiberiberi vitamine" of Funk (also known as the antineuritic factor or water-soluble B) might be a complex was suggested by the work of Emmett and Luros" on the basis of differences in susceptibility to heat destruction Smith and Hendrick" showed that autoclaved yeast, in which the antineuritic vitamin was destroyed,

TYPICAL MICROBIOLOGICAL DETERMINATIONS OF THE VITAMINS

ap	35	VIT	d daa ariika	LFICI	EVCY	DISLA	SLS		1
		Response References for Preparation of Media Measured and Delants of Tests	Turbidity Analyst, 74, 310(1949)	US Pharmacopera XIV	Acadity US Pharmacopeia XIV	J Biol Chem, 192, 181 (1952)	Turbidity Ind Eng Chem, Anal Ed, 15, 141 (1943), J Biol Chem 160, 1	(1) US Pharmacopera MV, 3rd Sup (2) 4 Assoc Off Agr Chem, 36,	Acidity Proc Soc Exp Biol Med, 56, 95
		Response	Turbidity	Acadıts	Aerdity	Aeidity	Turbidity	Acidity	Acidity
	Vılamın ncentration	at Maxi- mum Grouth	ml tube 0 04	0 20	4	0 20	0 04	0 0003	0 0015
	Vılamın Concentration	at Half Haxi- mum Grouth	ы рег 10 ml tube 0 02 0 04	90 0	0 1	0 00 0	10 0	0 00005 0 0002	0 0004
TYPIGAL MICROBIOLOGICAL LATERATION		Test Mecroorganisms		Lactobacillus caser	Lactobacillus arabinosus 17-5 (Arcc 8014)	Lactobacillus arabinosus 0 03 17-5 (Arcc 8014)	Saccharomyces carls- bergensis	Lactobacillus letchmanss 313 (Arcc 7830)	Lactobacillus arabinosus
TYPICAL M.	Extraction and Ilydrolytic Procedure		Heat 30 mm 100° C 0 1 Lactobarellus fermentum N H ₅ SO, and digest 79 (Arcc 9833) with pepson and "Taka- Dustas".	Autoclave 30 min, 15 lb, Loctobacillus caser 01 N HCl	Autoclave 30 min, 15 lb., Lactobacillus arabinosus 0 1 10 N H ₂ SO ₄	Digest with intestinal phosphatase and liver enzyme	Autoclave 60 mm, 20 Ib, Saccharomyces carls- 0 055 N or 2N II ₂ SO ₄ bergensts	(1) Aqueous (USP) (2) 15 mm, 15 lb, pH 4 5 with fresh NaHSO ₁	Autoclare 60 mm, 15 lb, Laclobarillus arabinosus 0 0004 0 0015 6 N H SO,
		Vıtamın	Тылипе	Riboflavin	Nisein	Pantothenic acid	Vitamin B.	Vitamin B ₁₄	Biotin

Typical Michemological Detynamations of the Vitamins-(Confinued)

	The state of the s		Vita	Vitanin Concentration		
1.1/217111	F straction an l Hydrolytie Proce l cre	Test Ureroorganismis	at Half Vari	at Vaxi- rum Grouth	Response Meast red	Response (References for Preparation of Vitalia Vens red
o pouge	Drest by specific ontymes I actol neillus cases	I actol acultus cases	0 0002 0 002	0 002	Actibity	Act bly Arch Blochem 7, 287 (1015)
minobengoid	from chicken princreas	from chicken punceess (Art.c 71f))	0 015	0 01	Weight	Weight J Biol Chem 148, 281 (1913) modi
er I slimo	1f,SO, Autoclave 120 min 15 lb	(Arce 4273) Chobneless mutant of 6* Neurospora crassa No	•	0£	Weight	J Biol Chem 150, 325 (1913), Scrence, 101, 671 (1945)
sitol	Reflux 0 hrs 120° C 27 Saccharomyces carls- N HCl	34186 (ATCG 0277) Saccharomyces carls- bergensis (ATCC 9080)	3 0	8 0	Turbidity	Turbidity J Bact, 47, 131 (1944), Ind Eng Chem, And Fil, 15, 111 (1913)
						election measured to include pyrical
1						

Per 25 ml

still possessed supplementary value for rats receiving adequate amounts of rate min B from costs or in the form of a vitamin B picrate. Goldherger 46 in 1926, recognized that rats required not only the antineuritie vitamin ("B sensu stricto") but also another factor which he called P-P (nellagrapreventive) In 1927 the Accessory Factors Committee of the British Medical Research Council recommended the symbols B, and B, for the heat-labile (antineuritic) and the heat-stable components, respectively (For a time vitamin B. was called G in the United States) The erroneous helief that the latter was identical with the pellagra-preventive factor was corrected when distinction was drawn between human pellagra and canine blacktongue, on the one hand, and so-called rat pellagra on the other Retween the time that Jansen and Donath. 47 in 1927, isolated the crystalline anti-heriberi vitamin from rice bran and R. R. Williams 48 synthesized it and gave it the name this mine in 1936, evidence accumulated for the inereasing complexity of the vitamin B group

Kuhn, Gyorgy, and Wagner-Jauregg 19 isolated the fluorescent com pound ovoflavin from egg white, designating it vitamin B. Their demonstration that this biologically active substance and similar flaving from milk and liver (lactoflayin and hepatoflayin) were identical, containing a riboso group attached to an isoalloyazine ring, caused them to adopt the

common name riboflavia for vitamin B.

Evidence for the existence of additional growth and antidermatitis factors for rats, pigeons, and chieks prompted the continued search for new vitamins in the B group, which in the years sinco 1927 has culminated in the identification and synthesis of at least ten additional vitamins. From the standpoint of human nutrition, one of the most significant is nicotinic acid (now called niacin), which was identified by Elvehiem, Madden Strong, and Woolley, 50 in 1937 as the canine blacktongue factor This compound had been known for many years and was actually isolated from rice polishings by Tunk51 in his effort to isolate "vitamine B" The therapeutic trial of nicotinic acid in human pellagra was rewarded with prompt and dramatic success

Clinical experience with beriberi or pellagra has emphasized that these diseases are often associated with deficiencies of other nutrients than thiamine or macin alone Therapeutie administration of individual crys talline vitamins often reveals the presence of underlying deficiencies which respond to supplementary dosage with other vitamins of the B group or with rich sources of the vitamin B complex such as yeast or liver In this connection it is significant that the relative proportions of the various members of the vitamin B group as found in natural sources, both plant and animal, are quite variable, for example, seeds and legumes are relatively higher in thiamine than in riboflavin, whereas in milk and leafy

⁴⁶ Goldberger Wheeler Lallie and Rogers US Pub Health Repts 41 297 (1926) Gold berger and Lillie Ibid 41 1925 (1926)

4 Jansen and Donath Mededed Dienst Volksgezondheid in Vederland Indië 16 186

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Will ams J Am Chem Soc 58 1063 (1936)
 Kuhn G, Orgy and Wagner-Jauregg Ber 66 317 576 (1933)
 Flechjem Madden Strong and Woolley J Am Chem Soc 59 1767 (1937)
 Funk J Physiol 43 395 (1911) 46 173 (1913)

vegetables the reverse is true Processing by heat, exposure to light, or other destructive influences effect still further variations in the quantita

tive relationship of the B vitamins

Bacteria present in the mammalian intestinal tract are capable of syn thesizing various vitamins of the B group, noteworthy among them being biotin, folic acid, and pantotheme acid Tactors which influence such bacterial growth-eg, nutrients in the diet of the host or bacteriostatic medication (like the sulfonamides)-may depress intestinal synthesis sufficiently to induce secondary deficiencies

The use of microbiological assay procedures has aided materially in the discovery of additional factors in the vitamin B group which are essential for the nutrition of certain microorganisms or laboratory animals but whose role as vitamins for man is not yet established. Among these factors are p-aminobenzoue acid, thioctic (hpoic) acid, L citrovorum factor, L bulgaricus factor, and vitamin Br (the latter required by the mealworm Tenebrio molitor) Examples of microbiological assays are given under Riboflavin, Pantothenic Acid, Macin, Vitamin Be, and Pteroylglutamic Acid In principle these methods are identical with the procedures for ammo acid assay described in Chapter 33 and illustrated on p 1064 A table outhning the conditions for extraction, hydrolysis, and optimal concentration for microbiological vitamin assays by selected methods is given on p 1131

THIAMINE

The term "deficiency disease" was first applied to a condition known as beriberi, which was common in southeastern Asia and the islands of the Pacific Ocean The similarity in pathology between this disease and poly neuntis observed in fowls restricted to a diet of polished rice, both of which could be cured by feeding the "silverskin" (1 e , pericarp and germ) of the grain, prompted Eijkman in the Dutch East Indies, to investigate the subject from the nutritional standpoint Eighman produced beriben in fowls on a dict of polished rice and prevented the syndrome by dietar) means His classic studies, reported in 1896-1897, paved the way for a host of other investigations which culminated in the formulation of a vitamin hypothesis Besides those Oriental countries where polished rice is the main article of diet, other places in which beriber has been observed are prisons or asylums localities where the diet is apt to he restricted or faulty, and war-stricken countries

After numerous attempts to isolate the vitamin by methods involving silver or lead precipitation crystallization of a picrate, adsorption, etc. the vitamin was finally crystallized from rice polishings by Jansen and Donath in 1926 in the very laboratories in which the original studies were conducted by Eijkman, who was able after thirty years to confirm the antincurrite activity of the crystalline vitamin hydrochloride A decade later the structure and synthesis of the vitamin were announced by R R Williams and his co-workers

Clinical cases of thiamine deficiency in the Western Hemisphere are rare today Subclinical cases of deficiency are probably more prevalent arising from a limited intake of the vitamin or from increased requirements such as are observed in pregnancy or lactation. The neuritis observed in eases of chronic alcoholism has been claimed to result from diminished intake of food as well as insufficient thiamine to metabolize the alcohol consumed.

Physiological and Clinical Aspects of Thiamine, 52 Thiamine is essential for the growth and metabolism of all animals as well as of many plants and microorganisms. Deficiencies are characterized by a variety of

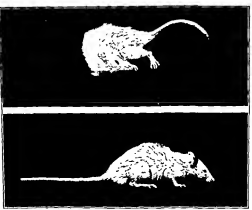


Fig. 276. Effect of Thiamine Deficiency and Subsequent Change to an Adequate Diet.

The spastic paralysis shown in the upper figure was cured (below) about 24 hours after addition of thismine.

Smith and Munsell: U.S. Dept. Agr. Circ. No. 84, 1929, Courtesy, The Bureau of Home Economics, U.S. Department of Agriculture.

symptoms and clinically are often complicated by the effects of lack of other nutrients.

In thiamine deficiency (avitaminosis B₁) a form of peripheral neuritis is manifested affecting both the sensory and motor nerves. During the early stages neuralgia and cramps of the calf muscles are common; as the condition advances the thigh muscles become weak and toe- and footdrop develop along with hypesthesia. The acute disease, beriberi, may be of the dry type, in which cachexia, aumbness, and paralysis are the primary symptoms, or of the wet type, associated with marked ("pitting") edema and paresthesia of the extremities. In animals the symptoms are

⁴³ Cowgill: "The Physiology of Vitamin B₄," in The Vitamins, Chicago, American Medical Association, 1939; Vedder: "The Pathology of Beriberi," bida; Straues; "The Therapeutic Use of Vitamin B in Polymeuritia and Cardiou ascular Conditions," bid

loss of muscular coordination, spastic movements, retraction of the head (opisthotonos), and paralysis (see Fig. 276). Recovery from the symptoms of polyneuritis is very rapid when thinmine is administered, especially by injection

Debility and progressive decline in weight are observed early in vitamin B₁ deficiency as a result of anorexia (loss of appetite), one of the most striking symptoms of this deficiency disease. The loss of appetite, which is the main cause of growth failure, has been attributed to a general systemic disturbance rather than to diminished secretory functions of the digestive glands or to reduced gastric motility, although diminished motor function of the gastrointestinal tract is observed chineally in thiamine deficiency.

The rapid restoration of appetite following thiamine supplementation of avitaminotic subjects has been regarded as indicating a specific appetite-stimulating function of the vitamin. The anorexin of thiminic deficiency, however, is more probably explained by the impaired carbohydrate metabolism of cellular tissue in general. Attempts to explain the loss of appetite on the basis of lowered secretory or motor function or diminished basal metabolic rate have not succeeded.

In polyneuritis pathological changes in the tissues and organs of the body include cardiovascular and neural disturbances and atrophy of the endocrine glands and other vital organs, but hypertrophy of the adrenals Prolonged deficiency results in cardiac failure and death. An outline of the clinical symptoms of thiamine deficiency is given in the American Medical Association syllabus on p 1289.

Degeneration of the myelin sheaths of peripheral nerves and also of the ganglion cells of the hrain and spinal cord is produced in experimental polyneuritis, but since similar findings are observed in atarvation, even when the supply of thismine is adequate, it is preferable to regard the primary neurological effect of the avitaminosis as a functional defect concerned with the physiology of the neurons. The rapid recovery from neurological symptoms which follows thismine therapy supports this view.

In 1937 Lohmann and Schuster** isolated from yeast a crystalline coenzyme, coearboxylase, the pyrophosphate ester of thiamme. When this compound is bound to a specific protein (apoenzyme) from yeast, and a magnesium ion, an active enzyme (carboxylase) is formed which catalyzes the decomposition of pyruvie acid to acetaldehyde and carbon diovde. Unphosphorylated thiamine bas no cocarboxylase activity, although it can stimulate the action of cocarboxylase in alkali-washed yeast preparations. The pyrinidine portion of the vitamin with an intact amino group has similar properties. Substitution of the magnesium by other divalent ions or combination of the eccarboxylase with different specific apoenzymes may result in more than one form of the enzyme and in somewhat diminished biological activity.

Free thiamine is phosphorylated in ruo to cocarboxylase (or diphos-

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⁵¹ Wolbach J Am Med Assoc, 168, 7 (1937).

¹⁴ Lohmann and Schuster Naturoussenschaften, 25, 26 (1937).

phothiamine) by hiver and kidney tissue and to some extent by muscle and brain. The phosphorylating agent appears to be adenosine-5'-triphosphate (ATP).

In animal tissues, cocarboxylase plays a role in various reactions involving principally the decarboxylation of pyrivic and other keto acids In the hrain, cocarboxylase participates in the anaerobic dismutation of pyrivate to lactate and acetate, and their subsequent oxidation to carbon dioxide and water. In liver and other tissue cells, cocarboxylase is involved in the conversion of pyrivate to oxidacetate which combines oxidatively and irreversibly with another molecule of pyrivate to enter the tricarboxyla each cycle. The oxidative decarboxylation of ketoglitarate to succinate and of pyrivate to acetoacetate, and the conversion of pyrivate to the acetyl group of acetyl-coenzyme A (see p. 1188), are additional examples of reactions found to be mediated by cocarboxylase.

It is not an exaggeration therefore to say that thinmine occupies a key position in at least the terminal stages of carbohydrate metabolism. In advanced thinmine deficiency both pyruvate and lactate accumulate in the blood whereas glycogen increases in the liver and beart muscle. Rats subsisting on diets of glucose or casein alone survive twice as long when the mine is added.

Diphosphothiamine may be involved in the synthesis of acetylchohno and in the control of its hydrolysis in ino. The activity of choline esteraso of serum is strongly inhihited he coerbovilsse.

The average American diet has been estimated to contain 13 mg of thiamine per day. Prior to bread and flour enrichment, the value was 08 mg Moderate thiamine deficiency may be the root of many ill defined nutritional disturbances, especially when anorevas is a symptom. Such cases respond readily to the therapeutie test, i.e., supplementation of the diet. The thiamine requirement is a function of sev. body weight, musicular activity, and other conditions. Diet, especially the intake of nonfat calories is an important factor. The recommended daily allowances of thiamine are shown in the table on p. 1108. The remarkable tolerance for thiamine is indicated by the fact that 500 mg have been taken daily for a month by normal people, without any objective symptoms. It is of interest in this connection that peripheral neutrits in chronic alcoholism has been attacked therapeutically on the hypothesis that the condition is due partly to a disturbance of the vitamin B₁ nonfat calorie ratio.

It has been shown that lactating rats require an additional quota of antineurity vitamin, in the absence of which the mother will neglect or destroy the litter. The vitamin may be supplied directly to the young or indirectly through the mother's mil. 45

Thiamine and its compounds are the only naturally occurring substances having vitamin B₁ activity for the higher species. Certain synthetic analogs have less biological potency, e.g. the compound having a methyl group in the 6-, rather than in the 2 position in the pyrimidine nucleus, the 2-ethyl, instead of the 2 methyl, compound is active for

¹¹ Sure J Biol Chem. 76 685 (1928) Sure and Walker J Biol Chem 91 69 (Macy Outhouse Graham and Long J Biol Chem 73 189 (1927)

Phycomyces; thiamine disulfide, a possible intermediate in the metabolism of thiamine, obtainable by mild oxidation of the vitamin, has full activity

Thiamine may be inactivated both in silro and in tito by an enzyme, thiaminase, present in certain fish and shellfish (see p. 1281). The vitamin is hydrolyzed to 2-methyl-6-amino-5-hydroxymethyl pyrimidine and 4-methyl-5-hydrovyethylthiazole.

Pyrithiamine, a mixture of derivatives which include the true pyridine analog of thiamine (i.e., 2-methyl-3-8-hydroxyethylpyridine in place of the thiazole moiety), and oxythiamine, in which oxygen replaces the sul-

fur atom, are powerful thiamine antagonists (see Chapter 36).

Storage and Synthesis of Thlamine. The available evidence seems to indicate little capacity on the part of the animal body to store thismine. It occurs both free and phosphorylated, somewhat higher concentrations being present in the heart, liver, and kidneys than in muscles and brain. It is possible to increase the thiamine content of the tissues by dictary means; however, their storage capacity is so limited that even under these circumstances only a few weeks' reserve can be maintained It is thus important that the daily diet melude an adequate supply of thiamino

Whereas certain microorganisms can synthesize the vitamin, others require an external source; some utilize the thiazole and pyrimidine portions-e.g , Phycomyces blalesleanus-whereas others can synthesize one moiety and combine it with the other obtained from an external source. Thiamino is synthesized by all higher plants, though to only a limited extent in the dark.

Though all animals require thiamine, sheep and eattle do not need an external supply since the vitamin is synthesized in sufficient amount by bacteria in the rumen. Though thiamine is known to be synthesized by intestinal flora in rats and humans, the nutritional importance of this contribution is not known Difficulties encountered in the production of experimental thiamme deficiency in humans subsisting on a diet low in thiamine have been attributed to the intestinal synthesis

Thiamine and eocarboxylase have the same biological activity for higher animals Thamme can be phosphorylated by live yeast, by dried yeast, or by a phosphatase obtainable in the presence of hexosediphosphate, adenosine triphosphate, and a specific protein The vitamin may also be phosphorylated by certain bacteria. Liver and kidney, and to some

extent muscle and brain, can perform the same function Distribution of Thiamine. 55 This vitamin is essentially of plant origin There is no proof of its synthesis by the animal body proper although it is synthesized to some extent by bacteria in the intestinal tract. Its presence in such products as milk, eggs, and liver depends on the dietary supply of the animal Of the various meats used for human consumption, pork muscle is the richest in thiamine The ability to synthesize thiamine is possessed by plants and by lower forms of hie, e g, bacteria, molds, and yeasts The latter are a potent natural source of this factor, as well as of other components of the vitamin B complex.

⁴⁴ For thiamine values in foods see Appendix III.

Thiamine is widely distributed in the plant world, seeds, leaves, roots, stems, and fruits being recognized sources. In cereal grains as a class, thiamine is found in linghest concentration in the germ or embryo, less in the bran, and least in the endosperm. The amount of thiamine in a milled grain such as flour depends on the proportion of each part of the grain present. Leaves and grasses (e.g., spinach, alfalfa, timothy liny, etc.) are rich sources of thiamine as are also beaus, nuts, fruits, milk and egg yolk.

Vitamin B₁ may occur in natural materials as free thiamine, as a protein complex, as pyrophosphore acid complex (cocarhoxylase), and as a phosphorus-protein complex In milk, the vitamin is present both as free thiamine and as the vitamin protein complex In hrain and liver, it is found as cocarhoxylase and as its protein complex In wheat, thiamine occurs in the free form In blood, it is present as the pyrophosphate in the cells and as free vitamin in the serum. Thiamine is excreted in the free form in the urine.

Chemistry of Thiamine. The efforts of many investigators to isolate pure crystalline thiamine from natural sources culminated in the work of Jansen and Donath (loc cit) They adsorbed an acid extract of rice polishings on clay, cluted with Ba(OH), and alternately precipitated (first with silver nitrate, then with phosphotungstic or silicotungstic acid) and decomposed with acid, finally obtaining a platinic chloride precipitate from which a crystalline thiamine hydrochloride was prepared. The procedure yielded about 1 part of the vitamin from 3,000,000 parts of rice polishings

Analysis of the crystalline vitamin led to its synthesis by Williams and his co-workers in 1936, ⁵⁷ which was soon followed by large-scale commercial production Thiaminc chloride is 3-(4'-amino-2'-methylpy rimidyl-5'-methyl)-5-\(\theta\)-thought drove the 1-l-methylthiazolium chloride. Its hydrochloride, commonly called thiamine chloride, is

Thiamine chloride hydrochloride (C₁₂H₁₁_SOCI HCl)

Thamme hydrochlorde is a white crystalline solid. The crystals are stable at 100° C for 24 hours but decompose when heated to the melting point of 249° C. The compound is hygroscopic and dissolves in water to form an acid solution which is optically mactive. One grum dissolves in 1 ml of water, in 18 ml of glycerol, in 100 ml of 95 per cent alcohol, or in 315 ml of absolute alcohol. It is insoluble in ether, acctone, chloroform, and benzene. The vitamin has a characteristic yeasty odor and taste.

Williams J in Clem Soc 58 1063 (1936) Cline Williams and Linkelstein J Am Chem Soc 99, 1052 (1937) Williams Ind Eng Chem 29 980 (1937) The synthesis of thrumno was also accomplished at about the same time by Todd and Bergel (J Chem Soc, p 364 (1937)) and by Andersag and Westphal (Ber \sim 2035 (1937))

assay of urme by the throchrome method. In the analysis of materials that yield clear solutions and low blanks, preliminary tests may justify omission of the adsorption-elution step (in both the standard and unknown) Examples of such materials are certain pharmaceutical preparations, enriched flour, meat extracts, and milk

In the colormetric procedure, the cluate from the zeolite is first added to a solution of phenol in alcohol to increase the sensitivity of the test The solution is then made alkaline and diazotized p-aminoacetophenone is added The mixture is allowed to stand while the red dye develops. The latter is then extracted selectively by xylene which leaves behind in the aqueous phase dyes produced by the coupling of the reagent with phenol, phosphorylated thiamine (if present), acid base indicators, and other interfering compounds Results obtained by the colorimetric procedure agree well with those obtained by biological assay

Since many microorganisms require an external supply of thiamine for growth, microbiological methods may be employed for the determination of this vitamin Caution should be exercised in the choice of microorganisms, however, since some can utilize portions of the thiamine molecule, or degradation products of the vitamin which are not biologically active for higher organisms Successful microbiological assays have been con ducted employing Lactobacellus fermentum 36, the growth of which, under the conditions of the test, is not stimulated by the pyrimidine or this zole portions of the thiamine molecule, either alone or together Results obtained by yeast fermentation methods are erroneous if the pyrimidine portion is present, since it stimulates yeast fermentation Correction for this interference may be made by conducting a test on a sulfite-treated sample in which the thiamine is inactivated 62 63

Cocarboxylase may be determined specifically and directly by measurement of the carbon dioxide produced by decarboxylation of pyruvic

acid by yeast cells previously washed free of cocarboxylase "

Human thiamine deficiency, both chinical and subchinical, may be diagnosed by several biochemical procedures. These my oly c measurement of the concentration of thiamine, cocarboxy lase or pyruvic acid in the blood or urine Since the levels of these compounds may fluctuate widely depending on the dietary intake for several days immediately preceding the test, a more reliable diagnosis is made when the response to a test dose of the vitamin is measured A simple and reliable diagnostic test 65 68 18volves colonmetric measurement (see p. 1144) of the urinary exerction of thirmine in the 4-hour period following intramuscular injection of 0.35 mg of the vitamin per square meter of body surface, 12 hours after the last meal Normal individuals exerete more and deficient subjects less than 50 ng of thamme during the 4-hour period

CHEMICAL METHODS

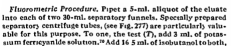
Modification of the Fluorometric Method of Hennessy and Cerecedo: frinciple. The thiochrome method for the determination of thiamine involves extraction of the vitamin, dephosphorylation, purification by adsorption-clution, and oxidation. The fluorescent thiochrome is extracted with isobutanol and determined fluorometrically. For a general discussion of fluorometry, see Chapter 23.

Procedure: Preparation of Sample. Extract a sample containing not more than 20. of solids by refluxing for 0.5 hour with 75 ml. of 0.1 N sulfuric acid.

Cool to 40° C. If the material is rich in protein, add 0.5 g, of pepsin and incubate over night at 37° C. Bring to 100° C. and then cool. Add 10 ml. of 1.8 M sodium acetate and adjust the pH to 4.5 If necessary. Add 0.5 g, of "Taka-Diastase" and incubate overnight at 37° C., or 3 hours at 45-50° C. Cool. Dilute to 100 ml. with distilled water, and filter.

Pass an allquot of the filtrate containing approximately 3 5 µg, of thiamine through a base-exchange tube* containing a column of activated zeolite.* Wash the column twice with 20-ml. portions of distilled water. Elute the vitamin with 25 ml. of 25 per cent potassium chloride in 0.1 N hydrochloric acid. Mix the cluate.

Pass a standard solution containing 5 rg. of thiamine in acidulated water (pH 4 5) through a separate column, and elute as above.





Courtesy, Scientific Glass Apparatus Co Bloomfield,

then 3 ml of 15 per cent sodium hydroxide to the other, the blank (B). Shake the funnels for 1.5 minutes, centrifuge, and discard the aqueous phases. Clarif, the isobutanol layers by shaking with approximately 1g. of anhydrous sodium sulfate, transfer to cuvettes, and read in a fluorometer equipped with appropriate filters. It use an aqueous solution of quinine sulfate containing 0.2 we, ner ml. in 0.1 N sulfuric acut to check the setting of the instrument.

⁴⁷ Hennessy and Cerecedo J Am Chem Soc, 6t, 179 (1939), Hennessy Ind Eng Chem, Anal Ed, 13, 216 (1941)

⁴⁵ The base-exchange tube contains a reservoir at the upper end having a capacity of approximately 30 ml, followed by a tube of 5 to 6 mm internal diameter approximately 14 cm long. At the lower end is a capillary of such diameter that when the tube is charged, the rate of flow will be supportunited.

the rate of flow will be approximately 1 ral per minute

"Approximately 50 mesh "Decaleo" Activate the zeolite by stirring with four 10-volume
portions of 3 per cent acetic acid for 10 minutes each. Between the second and third acid
wash treat for 15 minutes with 5 volumes of 25 per cent potassium chloride solution. Wash
the zeolite with water till free of chloride then with alcohol, and ether Dry in air and store
in a sealed bottle Place a pledget of glass wood over the upper end of the capillary, and pour
a water suspension of 2 g of the activated zeolute into the tube

¹⁶ Dissolve 30 mg of potassium ferriev anade in 100 ml of 15 per cent sodium hydroxide. ¹¹ The fluorometer should be equipped with an ultraviolet light source and a filter with a transmission peak at 3700 Å. Between the glass cuvette and the photocell insert a secondary filter with a transmission peak at 4600 Å. Since thiochrome is unstable to light, make the measurements rapidly and in a sexuidar room.

CALCULATION Calculate the thiamine content of the sample employing the formula

$$\frac{T_u - B_u}{T_s - B_s} \times \frac{1}{5} \times \frac{25}{A} \times \frac{100}{G} = \mu g$$
 thiamine per g of sample

 $T_{\rm w}$ equals the galvanometer reading obtained with the isobutanol extract of the unknown treated with ferneyanide, $T_{\rm w}$ is the corresponding value obtained with the standard $B_{\rm w}$ and $B_{\rm s}$ are the galvanometer readings obtained in the blank tests conducted on the unknown and standard respectively, A is the volume of the unknown solution passed through the column and G is the weight of sample taken for analysis

2 Colorimetric Method of Hochberg, Melnick, and Oser in Principle The red pigment formed in the reaction between thismine and diazotized p-animoacetophenone is measured in a photoelectric colorimeter. The method involves the preparation of a clear extract of the free utamin, adsorption on and elution from a reolite column reaction with diazotized p-aminoacetophenone extraction of the red pigment with xylene and comparison with a standard similarly treated. The method is applicable to the determination of thismine in urine. Here the extraction and enzyme hydrolysis are omitted and the urine samples after adjustment of the pH to 4.5 are passed directly through the reolite column.

The use of large samples for analysis makes the colorimetric method sufficiently sensitive for the determination of thismine in most products. However, it is not recommended for the assay of materials rich in protein but low in thismine. Because large samples are taken for analysis high concentrations of adsorbable amino acids are released which interfere with the retention of thismine on the scolite column.

through the outside jacket, and pour 30 ml of water on the column Allow to heat for one-half minute, then draw through with full suction to wash and heat the zeolite adsorbate Elute the thlamine Immediately by passing 10 mi of 25 per cent potassium chloride solution in 0 1 N hydrochioric acid down the wall of the hot condenser Collect at the rate of approximately 1 drop per 2 seconds, drawing through the final few drops by suction Wash the zeolite column with 200 ml of distilled water under full suction and with the steam on Cool the column by running the last 50 mi of wash through with the steam turned off The apparatus is then ready for the next extract

Pass a standard splution containing 30 µg of thiamine in 50 mi of acidu lated water at nH 45 through the column in the same manner as the test overget

Colorsmetric Procedure Transfer the cluate to a 100-ml centrifuce tube Add in mi of alcohol phenol reagent," previously poured into the receiving tube as a wash, then 2 drops of a 1 per cent alcoholic solution of thymol blue Bring all samples to this point before proceeding Handling one tube at a time, add 2 N sodium hydroxide dropwise, with constant stirring, till the first distinct blue color is produced, then immediately add 25 ml of freshly prepared thiamine reagent " Ailow to stand at least 2 hours, or preferably overnight, at room temperature Add 5 to 15 ml of xylene and shake vicorously for 3 minutes Centrifuge Transfer the xylene layer by means of a Il tinned pipet to a Nessler tube, a visual colorimeter cup, or the absorption cell of a photoelectric colorimeter (520 mm filter), and evaluate versus the noituios brahases

CALCITATION The color intensity is linear in the range recommended. Calculate the thiamine content of the sample using the formula

$$\frac{U}{S} \times 30 \times \frac{200}{G} = \mu g$$
 thusmine per g of sample

U/S is the ratio of the concentration of red pigment in the unknown to that in the standard (the ratio of the photometric densities when a photoelectric colorimeter is employed) G is the weight of the sample in grams

BIOLOGICAL METHODS

The symptoms of thiamine deficiency which have been adapted to biological assay are polyneuritis and anorexia. Pigeons, 77 chicks 78 and

⁷⁶ Dissolve 3 9 g of phenol in 500 ml of 95 per cent alcohol Store in an amber glass bottle Dissolve 0 635 g of p-ammoscetophenone in 9 ml of concentrated hydrochloric acid and dilute to 100 ml with distilled water Dissolve 22 5 g of sodium nitrite in distilled water and dilute to 500 ml Dissolve 20 g of sod um hydroxide and 28 8 g of and um hydroxide honate in d stilled water and dilute to 1000 ml

Diazonium Salt Solution Pipet 5 ml of p-aminoacetophenone solution into a 50-ml eylinder surrounded with chopped ice and water and provided with a stirrer. Add 5 ml. of sod um nitrite solution slowly and stir for 10 minutes. Add an additional 20 ml. of nitrite solution slowly with stirring and keep in the ice bath for 30 minutes longer. Store at a tem perature below 5° C and use the solution on the same day it is prepared

Thumine Reagent Add 10 mi of the diazonium salt solution to 137 ml of sodium hydrox

ide-bicarbonate colution with vigorous stirring. Allow to stand till the initial pink coloration changes to pale yellow (5-20 minutes) then use immediately

changes to paic yethow (o-20 minutes) then use immediately in Kinnerske; Peters and Reader Backets J 22 276 (1929) Coward Burn Ling and Morgan ibid 17 1"19 (1933) Waterman and Ammerman J Nutrition 10 161 (1935) Carter and O Brien Bochem J 31 2"64 (1937)

14 Arnold and Elvehjem J Aufrition 15 403 (1937)

rats79 are employed in the methods hased on cure or prevention of poly neuritis, which have the advantage of specificity but are not as adaptable to quantitative treatment as are growth methods Curative methods are based on the duration of the cure of head retraction in birds or of paralytic convulsions in rats, resulting from feeding graded doses of the test material The difficulty arises in determining precisely when the symptoms have advanced sufficiently to administer the test dose and when "cure" is effected, the question being one of hours rather than days and subject to considerable personal judgment. The pigeon-curative method has strong defenders, particularly among English workers Before the differentiation of vitamin B was established, most of the studies on the antineuritic properties of the vitamin B complex were conducted on p geons, because rats rarely exhibit typical symptoms of polyneuritis even though they cease to grow when deprived of this complex Polyneurities can be produced in rats by maintaining a state of partial deficience in order that the rats survive long enough for the symptoms to appear method hased on the growth response of rats to graded doses of thiamine has been adopted by the Association of Official Agricultural Chemists

In the complete absence of thiamine, wearling rats show a gain in weight for the first week or so, followed by a rapid decline and death in 25-40 days If life is prolonged on a submaintenance allowance of this mine, the characteristic picture of polyneuritis appears, including lack of appetite, torpidity, spastie movements of the head and legs, and, in more severe cases, head retraction and paralysis of the hind legs Lack of muccular coordination and a tendency to hold the head to one side and walk in circles are frequently manifested (See Fig. 276, p. 1135.)

After 4-7 weeks, evidences of polyneuritis are seen, including incoording nation, spasticity, and rolling movements Spinning the rat by its tail will usually evoke convulsive seizures When these become distinctive and consistent, the test foods or extracts are fed or injected in sufficient dosage to effect cure, the quantitative criterion being the minimum dose which will prevent a return of the symptoms for at least five days This method, as adapted by the U S Pharmacopera, is described in full on p 1148

The rat-growth method of Sherman and Chases has been the basis of many evaluations of the vitamin B₁ content of foods and, except for the composition of the basal diet is substantially the biological assay method

for thiamine tentatively adopted by the AOAC (see below)

Coprophagy must be prevented in assays for the B vitamins since bacterial synthesis in the intestinal tract may produce significant concen trations of vitamins Thus rats may grow on a B-free diet, this condition being known as "refection "si Partial digestion (dextrinization or gelatini zation) of the starch in the diet by boiling has been recommended to avoid refection

Rat Growth Assay for Thiamine "Principle Rats depleted of their thiamine reserves by means of a vitamin B_i free basal diet are fed graded supplements of thiamine to the form of the standard or test materials. The relative growth responses to the doses of assay material and of pure thiamine constitute the basis for evaluation

Procedure Health, rats not exceeding 28 days of age and weighing between 40 and 50 g are placed in individual cages with raised wire mesh floors (not less than 8 \times 8 mm mesh) Water and basai diet are fed ad libitum

BASAL THIAMINE DEFICIENT DIETS

Sterman and Chase		AOAC		
Caselo (thiamine-free)**	18	Casein (thiamine-free)*3	18	
Starch**	53	Sucrose	60	
Salt mixture*6	4	Salt maxinres	4	
Butterfat**	8	Liver extract	1	
Codhyer oilss	2	Codhver oil	2	
Autoclaved baker s 3 east 17	15	Autoclayed yeast*7	5	
•		Autociaved peanuts**	10	
		P3 ridovine	0 0002	

The rats are weighed at intervals not exceeding 3 days. After 10 (but before 30) days the rats are depleted, as evidenced by stationary or declining weight over any 5 day period. They are then assembled into groups of 8 or more, one group being maintained on the hasal diet as the negative control, the remainder receiving either graded doses of USP thiamine hydrochloride (0 1 ml = 30 μ g thiamine) fed directly into the mouth by means of a syringe with a hiunt edged 18 gauge needle or measured doses of the assay material corresponding in expected thiamine content to the amounts of standard fed to the reference groups. The intervals between doses should be constant for both reference and assay groups each dose is computed by multiplying the next lower dose by a constant factor (e.g., 15 or 2).

CALCULATION During the 4-week assay period we ghts are recorded at weekly in tervals. The average net gains of the reference groups are plotted and the average gains of the assay groups interpolated on this curve in terms of thiamine content for each dose level of the assay material.

²² Cited by Sherman and Smith The Vitamins 2d ed New York Reinhold Publishing Corp. 1931. Association of Office al Agricultural Chemists. Official and Tentative Methods of Anglusia of the Association of the All Publishers. D.C. 1959.

of Analysis of the Association of the Washington D C 1950

Thau ine-free eveni may be prepared as follows Str 400 g of case n with 2 liters of 00 per cent (by weight) shooled for ½ hour let stand 6½ hours filter with suction and wash with 1 liter of 00 per cent alcohol Repeat letting stand 18 hours and adding a final wash with 1 liter of 90 per cent alcohol Repeat letting stand 18 hours and adding a final wash with 1 liter of 90 per cent (by we gith) shooled Spread on trays to dr in air 1 vita in free casein is supplied by the Borden Co Bambridge N X and by General B ochemicals Chagrin Falls Oi to

[&]quot;See p 1146
"See USP salt m rture p 1269

[&]quot;In place of butterfat and codiver ol 10 per cent hydrogenated vegetable oil may be used togetler with "000 U.S P units of vitama A and 200 units of vitama D per 100 g of basal det added in the form of high potency fish liver oil or concentrate."

[&]quot; See footnote 93 p 1150 " See footnote 94 p 1150

If desired the data may be treated statistically according to the analysis of variance illustrated for vitamin A in Appendix VI

Results are preferably expressed in gravimetric units-eg, mg of thiamine per 100 g of assay material the International nr US Pharmacopeial unit is the potency of 0 003 mg (3 µg) of cry stalline thiamine hydrochloride

ASSAYS FOR THIAMINES

Biological Method of the U.S. Pharmacopeia XIV.

The biological assay, comprising the recording of observations of rats through out specified periods of their lives, while being maintained on specified dietary repmens, and the interpretation of such data, is as follows

PRELIMINARY PERIOD Throughout the preliminary period each rat shall be raised under the immediate supervision of, or according to directions specified by, the assayer Throughout the preliminary period the rats shall be maintained on a dietary regimen which shall provide for normal development in all respects, except that the thiamine hydrochloride intake may be restricted

DEPLETION PEPIOD A rat shall be suitable for the depletion period when the age of the rat does not exceed 30 days, and if the body weight of the rat does not exceed 50 g and if the animal manifests no evidence of injury or disease or anatomical abnormality which might hinder growth and development Throughout the depletion period each rat shall be provided with the thiamine hydrochloride test diet and water (USF) ad libitum, and during this period no other dictary supplement shall be available to the animal Throughout the depletion period and until the assay shall have been completed, the rats shall be kept in cages provided with a wire cloth bottom, each mesh of which shall be not less than 8 mm

Assay Preion A rat shall be suitable for the assay period, provided that the depletion period shall not have exceeded 75 days, and provided that the rat shall manufest evidence of thiamine hydrochloride deficiency characterized by acute polyneuritie Throughout the assay period each rat shall be kept in an individual cage and provided with the thiamine bydrochloride test dat, compounded from the same lots of ingredients and water (USP) ad libsium On the day beginning the assay period there shall be administered to each rat a single dose of the reference standard of such size that it will produce in individual animals a curative period of not less than 5 days and not more than 15 days. All of the rats used in any one assay shall receive the same quantity of the reference standard Each rat shall then be observed for the cure of and recurrence of polyneuritis and when polyneuritis reaches the same acute stage observed when the reference standard was administered, a single dose of the assay product shall be administered. The animals shall then be observed to determine if polyneuritis is cured, and if so observation shall be made of the duration of the period Each assay shall include successive administration of the reference standard and assay product to not less than eight rats. The assay product may be administered orally or parenterally, but in any one assay the reference standard shall be administered in the same manner as the assay product, and the quantity of the assay product administered to each rat shall be the same

Recomping or Data. On the day beginning the depletion period and at intervals of not more than 7 days during the depletion period a record shall be made of the body weight of each rat On about the twenty fifth day and each day thereafter for the remainder of the depletion period each rat shall be observed for symptoms of polyneurits. The following days in the contraction of the contrac polyneuritis. The following dates shall be recorded

1 The day on which the reference standard is administered.

[&]quot;Grateful acknowledgment for permission to reproduce these methods (U.S.P. XIV) is made to Dr Lloyd C Miller and the Board of Trustees of the U.S. Pharmscoperal Convention Inc.

- 2 The day on which cure of polyneuritis is observed following the administra-
 - 3 The day on which acute polyneuritis recurs and the assay product is fed
- 4 The day on which cure of polyneuritis is observed following the administration of the assay product
- 5 The day on which acute polyneuritis recurs after the administration of the assay product

THAMINE HYDROCHLORIDE OR NITAMIN B, POTENCY OF THE ASAY PRODUCT In letermining the thiamine hydrochloride potency of the assay product, the duration of the curative period following the administration of the reference standard and the assay product shall be considered. The dose of the assay product administered contains an amount of the thiamine hydrochloride equal to or greater than that contained in the dose of the reference standard administered if that quantity promotes in the assay animals a total curative period (the sum of the number of days of the curative period of each of the animals) equal to or greater than the total curative period produced by administration of the reference standard.

DEPOSITIONS As used herein, unless the context otherwise indicates, the term acule noluneurilis means that stage of thiamine hydrochloride deficiency in which the animal regains control of the voluntary muscles, as evidenced by standing or walking a few seconds after extreme muscular contraction, which has been induced by twirling the rat by its tail (the onset of acute polyneuritis is invariably accompanied by loss in hods weight) The term assay period means the interval in the life of a rat between the last day of the depletion period and the final observation following the administration of the assay product, the term areay product means a product under examination for its thiamine hydrochloride potency, the term curaire period is the interval of time between the administration of this mine hydrochloride and the subsequent regirrence of source polymenritis after a complete di appearance of polymeuritis symptoms, and the duration of the curative period is expressed as the number of days in that interval. the term cure of polymeuritis means the complete disappearance of polyneuritic symptoms and is invariably accompanied by increase in body weight, the term depletion period means the interval in the life of a rat during which its food intake is only the thinmine hydrochloride test diet and water (U.S.P.), the term preliminary period means the interval in the life of a rat prior to the depletion period, the term reference standard means the U.S.P. Thiamine Hydrochloride Reference Standard, the term thiamine hidrochloride test diet means a uniform mixture, which has not been compounded for more than 7 days, of the following food materials and in the proportions designated

Suggestions for Using the U.S.P. Thiamine Hydrochloride Peference Standard. Before preparing a solution of the Reference Standard dry it to constant weight in a

desiceator over phosphorus pentonide

Precautions to be Taken in the Preparation of Solutions Because of the hygroscopic nature of the completely desiceated U.S.P. Thismane Hidrochloride Reference Standard it is preferable to transfer the quantity required for a test to a small glass-stoppered weighing bottle in which it can then be weighed on a microbalance or an ordinary balance according to the number of tests for which it is to be used. Even without such precautions however exposure to air during weighing will not cause an increase in weight of more than about 0.6 per cent of the operations are completed within a minutes.

Ventral and all-aline solutions of thannies by drockloride are unstable and water acid solutions are readily infected by molds which mactivate the vatuum. Therefore stock solutions should be prepared using 20 per cent alcohol and containing sufficient by drockloric acid to make the solution approximately 0.002. A convenient strength for a stock solution is 0.5 mg. of thannie hydrochloride to each rol. These solutions are stable if stored at about 4.9

Solutions of suitable strength for animal docage (20 to 100 µg, per ml) must be made at least twice weekly from the stock solution by dilution with water. Such dilutions must be kept at a low temperature and examined dails for mold.

down all of the squeate from the walls of the reservoir. To keep air out of the adsorption column, a layer of liquid must be kept about the surface of the sile ite throughout the adsorption process and the tube may be prevented from draining by placing a rubber cap (falled with water to avoid inclusion of air) over the lower end of the capillary.

NEUTRAL POTASSIUM CILLORINE SOLUTION Dissolve 250 g of reagent potassium

ACID POTASSIUM CHLORIDE SOLUTION Add S 5 ml of reagent hydrochloric acid to

Somina Hydroxide Solution, 15 PER CENT Dissolve 15 g of sodium hydroxide

POTASSILM FERRICANTHE SOLUTION, I PER CENT Dissolve 1 g of reagent potassium ferricantide in sufficient water to make 100 ml. This solution must be freshly prepared on the day it is used

ONDELY G REACENT Trepare the solution by mixing 40 ml of the 1 per cent potas sum ferries and e solution with sufficient of the 15 per cent sodium by droude solution to male 100 ml. This solution should be used within 4 hours.

QUINNE SULFATE STOCK SOLUTION Quinne sulfate solution is used to govern the reproducibility of the fluorophotometer A stock solution of quinne sulfate is prepared by dissolving 10 mg of quinne sulfate in sufficient 0.1 A sulfatine acid to make 1000 ml. Present this solution in high resistant containers.

QUINIX SULFATE STANDARD SOLUTION Dilute 1 volume of the quimne sulfate stock solution with 30 volumes of 0.1 N sulfure and This solution finoresces to approximately the same degree as the throchrome obtained from 1 µg of thiamine hydro-chloride. Preserve this solution in held resistant containers

Thiamine Hyprochloride Stock Solution Weigh accurately not less than 25 mg of USP Thamine Hydrochloride Reference Standard which has been kept in a desiccator over phosphorus pentovide for at least 16 hours Since the Reference Standard is hygro-copic, precautions must be taken to avoid absorption of moisture Dissolve in 20 per cent alcohol adjusted to a pH of 3 5 to 4 3 with hydrochloric acid and make up to a volume of 1000 ml Store in a cool place in a well-closed, light-resistant container

THABLYE HYDROCHORIDE STAYDERO SOLUTION From a portion of the stock solution that has been warmed to room temperature, transfer to a 100-ml volumetric flask an aliquot contuning exactly 100 mg of thamme hydrochloride, and dilute to 100 ml with water adjusted to a pH of 3.5 to 4.3 with hydrochloric acid. Each ml of this solution contains 1 mg of thiamme hydrochloride.

Dilutions of this solution are treated in the same manner as that used in the *Prepara* ion of Assay Solution with respect to acid digestion, enzyme treatment, adsorption, and clution from the base exchange subsets.

PREPARTION OF ASSAY SOLUTION. The amount of material taken for the assay should be such that the ratio of the volume of 0 1 N sulfure acid used for the extraction to the quantity of sample is at least 15 to 1, and the content of thiumine equivalent to 30 to 100 kg of thiamine hydrochlonde. Place the necurately weighed quantity of the material to be assayed in 65 ml of 0 1 N sulfure acid contained in a 100-ml centrifuge tube and digest it on a steam both, with frequent musing, for 30 minutes, or heat in an autoclave at 121° to 123° for 15 minutes. The liquid raust remain acid during the digestion. If it is not distinctly acid to the thymol blue pH indicator, add sufficient dibited sulfure acid to make it acid. Cool, and adjust the pH to between 4 and 45 by the addition of 2 N sodium acetate solution, using bromocresol green pH indicator in conjunction with a spot plate. Add 5 ml of the enzyme solution, mix, and incubite at 45° to 50° for 3 hours. Cool, centrifuge the mixture until the supermixtual liquid to a 100-ml valumetric flask. Wash the residue by

centrifuging with 10 ml, then with 5 ml of 0 1 N sulfuric acid. Add the washings to the supernatant liquid and dilute to 100 ml, with water

Pass through the prepared base exchange tube an aliquot of the solution estimated to contain 5 to 10 µg of thiamine wash the tube with three 5-ml portions of boiling water taking care to prevent the surface of the hind from falling below the surface of the silicate.

Elute the thamme from the base exchange silicate by passing successively through the tube small portions of hot acid polassium chloride solution. Collect the first to to 20 ml of the liquid (eluate) in a glass-stoppered, 23-ml volumetric flass, cool and dilute to a volume of 25 ml with acid polassium ehloride solution. This constitutes the casay solution.

Oxidation of Thiamine to Thiochrome and Measurement of Fluorescente Determine the thiamine content of the oxidized assay solution by comparing the intensity of fluorescence of an extract of this solution exposed to ultra violet rays ranging from 350 mg to 400 mg with that of oxidized Thiamine Ilydrochloride Standard Solution The intensity of the fluorescence is proportional to the quantity of thiamine present and may be measured with the aid of various instruments

To oxidize thismine to thiochrome add to quantities of the assay solution and of the similarly treated Thiamine Hydrochloride Standard Solution con taining from 0 10 to 2 pg of thiamine, sufficient acid potassium chloride solu tion to produce a volume of 5 ml Then add, with mixing, 3 ml of oxidizing solution Within 2 minutes add 13 mi of isobutyl alcohol and shake vigor ously for at least 11/2 minutes Centrifuge the mixture at a low speed until \$ clear supernatant solution is obtained Measure in a fluorometer the in tensity of fluorescence of the isobutyl alcohol solution directly if clear, or, if cloudy, after shaking with 2 g of anhydrous sodium sulfate Compare with this the intensity of fluorescence produced after oxidation of the properly prepared Thiamine Hydrochloride Standard Solution Quinne Sulfate Standard Solution is used to govern the reproducibility of the instrument Correction must be made for fluorescence produced by substances other than thiamine by determining the Intensity of fluorescence of Thiamine llydro chloride Standard Solution, and assay solutions treated as described above hut with 15 per cent sodium hydroxide solution replacing the oxidizing readent

The Thiochrome Method is applicable to substances such as Thiamine llydrochloride, and a number of other products, but cannot be relied upon when certain interfering substances are present, in the latter case, use the Biological Method, p. 1148

RIBOFLAVIN

growth-promoting properties of the vitnmin on rats Riboflavin was synthegrand shortly thereafter \$5

The clinical syndrome of amhofinamosis was first described in 1938 ha Sehrell and Butler 99 Arrhoflavinosis occurs more frequently among infants and children 100 than among adults and since its prevention depends upon an adequate diet it occurs most often among low income grouns 101 It is one of the more common dietary deficiency diseases and is usually associated with lack of other vitamins, notably macin. It is characterized by the occurrence of cheilosis and glossitis

The naturally occurring flavins—lactoflavin, ovoflavin, hepatoflavin. and verdoflavin-isolated respectively from milk, eggs, liver, and grass.

are all chemically identical with rihoffax in

Physiological and Clinical Aspects of Riboflavin, Vitamin B. deficiency affects primarily the ectodermal tissues, producing lesions of the skin eye, and nervous system. One of the earliest symptoms is cheilosis. manifested at first by transverse fissures at the corners of the mouth, raw and scaly lips, and finally by many vertical, deep fissures. The tongue assumes a purplish or magenta tinge and glossitis (flattening of the papillae) is observed A schorrheic dermatitis occurs at the body folds, e.g. at the alge nast, and in the scrotal and vulvar regions. The ocular manifestations of riboflavin deficiency include dryness, hurning and itching, photophobia and lacrimation, and vascular invasion particularly at the scleral junction of the cornea Cataracts due to pigmentation and capillary invasion of the cornea are known to occur in animals on a riboflavindeficient diet. but it is uncertain to what extent human cataracts result from dietary causes Though these symptoms may not individually he specific, choilosis, glossitis, seborrheie dermatitis, and corneal vascularization constitute a group of signs which separately or in combination are observed in amboffavinosis The symptomatology of clinical riboffavin deficiency is outlined in the syllahus on p. 1291

Clinical tests of the riboflavin content of blood and urine have been employed for the diagnosis of riboflavin deficiency. Normal individuals have a blood level of 0.5 ug per ml and when subsisting on an adequate diet excrete 500 to 800 µg per day in the urine. The excretion and blood levels show diurnal variations depending upon the dietary intake A diagnostic procedure involves measurement of the urmary excretion for 4 hours following the intravenous injection of 1 mg of riboflavin 10° Blood rihoflavin may be determined by the microbiological procedure, employing Lactobacillus cases, or by the fluorometric metbod. In addition to riboflavin, the latter measures proflavin, a metabolite of the vitamin

Riboflavin is required by all animals and many microorganisms. A deficiency of the vitamin in young animals results in inhibition of growth terminated by death. In rats, the syndrome includes early atrophy of the

³⁵ Kuhn Rememund Wevgand and Strobelo Ber 68, 1765 (1935) Karrer and associ

ates Hilds chim acta 18 1435 (1935)

**Sebrell and Butler Pub Health Rep 53 2292 (1938) 54 2121 (1939)

**Sebrels Bean Vilter and Hinff Am J Mej Sec 200 197 (1940)

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¹⁰² Najiar and Holt Bell Johns Hopkers Hosp 69, 476 (1941)

testes, involution of the thymus, alopecia, cataract, and degeneration of the main peripheral nerve trunks. In fowls, riboflavin is essential for normal egg production and hat chability Peripheral nerve degeneration in chicks is responsible for "curled toe paralysis" In monkeys, riboffavin deficiency results in anemia and leukopenia

Riboflavin plavs an important role in many enzyme systems In 1932, Warburg and Christian isolated a yellow respiratory enzyme from yeast 16 This enzyme was shown to consist of a combination of riboflavin phosphate and a protein The riboffax in phosphate, a mononucleotide, could be separated from the protein (apoenzyme) by dialysis against a weak acid Neither fraction alone possessed enzyme activity, but the two could be recombined in neutral solution to produce the original enzyme The yellow enzyme of Warburg and Christian can participate in a series of enzyme reactions involved in the metabolism of carbohydrates. It is capable of transporting hydrogen from reduced coenzyme II, a macin coenzyme which attacks hexosemonophosphate, for example, regenerating that enzyme of first attack. The reduced vellow enzyme may be reoudized itself by molecular oxygen This series of reactions however, is extremely slow and is probably of no physiological significance. Two other enzymes cy tochrome C reductase¹⁶⁴ and L-amino acid oxidase, ¹⁶⁵ contain riboflayin phosphate The former transports by drogen from reduced coenzy me II to cy tochrome C at a rate which is sufficiently rapid to he physiologically important This suggests that riboflavin functions in the oxidation reduction metabolism of plant and animal tissues L-Amino acid oxidase catalyzes the exidation of 1-amino acids and of a bydroxy acids

Rihoflavin-adenine dinucleotide

Riboflavin also participates in euzyme reactions as a dinucleotide prosthetic group, consisting of riboflavin, two phosphoric acids, ribose, and adenine This coenzyme is found in vanthine oxidase, diaphorase, p-amino acid oxidase, a synthetic enzyme of Warburg and Christian, fumaric hydrogenase, liver aldchyde oxidase, and the Haas enzyme, Xanthine oxidase106 (Schardinger enzyme, aldehydrase) is found in liver and milk and catalyzes the oxidation of aliphatic and aromatic aldehydes, reduced coenzyme I (another niacin coenzyme) and of a number of purines (including xanthine and hypoxanthine). In the presence of air, the reduced enzyme transfers its hydrogen to ovygen, forming hydrogen perovide which inhibits further action. The accumulation of hydrogen perovide is prevented by the presence of catalase, an iron-porphyrin-protein enzyme which catalyzes the decomposition of hydrogen perovide to water and ovegen. The hydrogens of xanthine oxidase, like those of numerous other reduced enzymes, can be accepted also by methylene blue and by other oxidizing dyes. Two different diaphorases107 have been found in plants, animal tissues, and microorganisms. In the presence of suitable proteins, one of these oxidizes reduced coenzyme I and reduced coenzyme II. p-Amino acid oxidase108 found in animal tissues, notably kidney and liver, converts p-α-amino acids to α-keto acids. Different p-amino acids are acted upon at different rates by this enzyme Fumaric hydrogenase, another enzyme which contains riboflavin dinucleotide, catalyzes the reduction of fumaric acid in the presence of certain reduced dyes 109 to

¹⁰⁰ Schardinger: Z Untersuch Nuhr u Genussm , 5, 22 (1902).

¹⁰⁷ Dewan and Green. Biochem J , 31, 1069 (1937). Von Euler and Hellström: Z. physiol.

¹⁰ Krebs, Z. physiol Chem., 217, 191 (1933), Klan. Wochschr., 11, 1744 (1932); Biochem J., 23, 1620 (1935).

¹⁰⁰ Fischer and Eysenbach: Ann. Chem , 530, 99 (1937).

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Riboflavin-adenine dinucleotide

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¹⁰⁷ Dewan and Green Biochem J., 31, 1069 (1937). Von Euler and Hellström: Z. physiol. Chem., 252, 31 (1938)

¹⁰¹ Krebs: Z. physiol Chem., 217, 191 (1933), Klin. Wochschr., 11, 1744 (1932); Bioch. J., 29, 1620 (1935)

¹⁰⁰ Fischer and Eysenbach: Ann. Chem , 530, 99 (1937).

succinic acid. Aliphatic and aromatic aldebydes are oxidized by liver aldebyde ovidase,110 a nboffavin dinueleotide enzyme found in the liver

The nhoflavin enzymes called flavoproteins, may be dissociated more of mammals or less easily and the original enzyme reconstituted by recombining the mono- or dinucleotide with the original protein The synthetic enzyme of Warburg and Christian¹¹¹ consists of the dinucleotide added to the protein of the old yellow respirator, enzyme The new enzyme differs from the natural enzyme ohtained from yeast only in that it contains the dinucleotide in place of the mononucleotide as its prosthetic group, it has the same properties as the old yellow respirators enzyme. The Haas enzyme consists of the dinucleotide plus a protein from yeast and has the same activity as the old yellow enzyme

Free nhoflavin is phosphorylated in the intestines of higher animal. prohably by an enzymatic reaction in which a secretion of the adrenab plays an important part When iodoacetic acid (an inhibitor of many enzymatic reactions) is administered or when the adrenals are removed phosphor, lation does not take place and the vitamin is not absorbed from the intestinal tract Under these conditions, riboflavin is unable to support growth, whereas the phosphate is effective. Human blood cells are capable of synthesizing the dinucleotide from free rihoflavin both in rice and in ritro 112

Since flavoproteins are composed of both riboflavin nucleotides and specific proteins disturbances in metabolism may result from diminished levels of either constituent. Deficiency of dietary riboflavin results in a decreased concentration of p-amino acid ovidase in liver and kidner in and of xanthine oudase in liver its Both of these conditions may be prevented or improved by feeding nhofiavin. The p-amino acidoudse activity of liver from animals on diets low in rihoffavin may be restored by the in ritro addition of the nucleotide to the excised tiscue 118 On the other hand rats on a low protein diet do not retain dietary rihoflavin in the here a condition which may be prevented by the administration of methionine Likewise, liver slices from nhoflavin-deficient rats are unable to mactivate estradiol this ability is retained when methonine is fed These observations illustrate the interrelationship of riboflavin with protem metabolism and with endocrine function

Riboflavin is concerned with the regulators function of the hormoneinvolved in carbohy drate metabolism. The administration of the roune to rats causes the loss of liver gly cogen unless riboflavin and thiamine are fed simultaneously The administration of insulin is effective in deparcreatized dogs only when these two vitamins are fed Riboflavin has been employed successfully in the treatment of dark adaptation in certain cases where vitamin A was ineffective. The retina contains free riboflavin which is converted by light to a compound which is involved in stimula-

Riboflavin and its two nucleotides are the only naturally occurring compounds which have vitamin B₂ activity. However, several synthetic compounds have been prepared having approximately one-half of the activity of riboflavin. These vitamers lack a methyl group in the six or seven position, or have this group substituted by an ethyl group. Removal of both methyl groups results in the formation of a highly toxic compound. Substitution of an alkyl group in the three position destroys the vitamin activity of riboflavin.

The minimum requirement of riboflavin necessary to maintain tissue stores in man has been estimated as approximately 0.5 mg, per 1000 calories ¹¹⁸ The recommended dietary allowances of the Food and Nutrition Board of the National Research Council are about 0.6 to 0.7 mg, per 1000 calories as indicated in the table on p. 1108. These allowances are more liberal than minimal requirements in order to provide a margin of safety. Under normal conditions, with an adequate dietary supply of riboflavin, humans everete approximately one-third of their intake. When larger amounts are ingested, approximately half is evereted in the urine. Very large doses of riboflavin have been fed to dogs and rats with no evidence of toxicity

Of interest is the relation between riboflavin feeding and experimentally induced neoplasms. Ingestion by rats of 2-acetylaminofluorene, a carcinogen, has been shown to induce ariboflavinosis; 112 high intake of riboflavin (2 mg per 100 g. duet) inhibited the appearance of choline-induced

tumors 115

Storage and Synthesis of Riboflavin. Riboflavin is not stored to any considerable extent in animal organs, though higher concentrations are found in the liver and kidney than in other tissues. In plants the younger parts are richer than the older. Broccoli leaves contain twice as much riboflavin as the flower buds; the latter contain more than the stems. Ungerminated seeds, other than peas, contain little riboflavin but appreciable amounts are formed during germination.

Riboflavin is synthesized by most higher plants, by yeasts, and by some bacteria Tbough higher animals are unable to synthesize the vitamin themselves, it is produced to a variable extent by microorganisms in the intestinal tract. In the rat this process is influenced by the nature of the carbohydrate in the diet. The ingestion of dextrin and corn starch favors synthesis, whereas sucrose is ineffective. In ruminants, the contribution of riboflavin by bacteria in the rumen is so great that a dietary source of riboflavin is not necessary.

Riboflavin is synthesized commercially and the product is employed on a large scale in bread and flour enrichment and for pharmaceutical purposes. Natural concentrates prepared from whey, yeast, and the

¹¹⁶ Williams, Mason, Cusick, and Wilder: J. Nutration, 25, 361 (1943).

117 Wase and Allison: Proc. Soc. Fron. Red. Med. 73, 147 (1950).

Mses and Allison Proc Soc. Exp Biol. Med. 73, 147 (1950).
 Schnefer, Copeland, Salmon, and Bale Cancer Research, 10, 786 (1950).
 MicElroy and Goss; J. Nutriton, 20, 527 (1940).

anaerobic bacterial fermentation of distillers' slops are employed in

Distribution of Riboflavin. 100 Riboflavin is widely distributed in plant and animal trisues as free riboflavin as the phosphate, or as the adenine-dinucleotide phosphate, the latter accounting for 70-90 per cent of the total riboflavin in tissues. 101 The free vitamin is found, for example, in milk, urine, and retina Riboflavin nucleotides occur more or less firmly bound to proteins Excellent dietary sources of riboflavin are heart, liver, kidney, muscle, eggs, milk, green leaft viegetables, veast, and whole grain. The riboflavin content of the average American diet before and after large-scale enrichment of bread and flour was practiced was 14 and 16 mg per day, respectively.

Chemistry of Riboflavin. Riboflavin is 6,7-dimethyl-9-(p-1'-ribit') iscallovazine. It crystallizes in orange-yellow needles which melt at 252° C with decomposition Riboflavin is insoluble in acetone, bearene chloroform, and ether, and only sighthy soluble in water (12 mg per 100 ml at room temperature), but very soluble in alkaline solution. Derivatives more soluble in water, such as the monosuccinate, borate, phosphate and acetate, have been prepared for pharmaceutical use. The vitamin is adsorbed by fuller's earth in acid solution and by frankonit in neutral solution but not by tale, alumnum ovide, calcium carbonate kabin of kie-elguhr, it is eluted by 80 per cent acetone or by alkalies such as

Riboflavin may be converted to its mononucleotide, riboflavin phosphate, by treatment with phosphorus oxychloride, or by suspending it in a phosphate buffer with intestinal enithelial nowder.

Determination of Riboflavin, Chemical determinations of riboflavin are based upon colorimetric122 and fluorometric123 procedures. In the analysis of food materials, colored and fluorescent compounds often interfere not only with the colorimetric determination of rihoflavin hut also with the fluorometric procedure in which the activating and fluorescent light are quenched. Some interfering pigments may be destroyed in the chemical test by oxidation with potassium permanganate or by reduction with stannous chloride. Fluorescence measurement before and after reduction of rihoflavin by treatment with sodium hydrosulfite has been employed to correct for the fluorescence produced by interfering substances. Correction may be made for the quenching effect of foreign pigments or other factors by the internal standard procedure. One method for the chemical determination of riboflavin 124 involves the conversion of the vitamin by irradiation in alkaline solution to lumiflavin. The latter is extracted from acid solution by chloroform and then determined colorimetrically or fluorometrically.

The microbiological determination of riboflavin depending upon the growth stimulation of Lactobacillus case; is more generally applicable to the determination of the vitamin in foods than the chemical methods. The microbiological procedure is sensitive and specific, and has been adopted as an official method by the US Pharmacopeia. It has been found that the growth of Lactobacillus case; in the presence of riboflavin is stimulated by suspended particles such as fat and starch. It is therefore essential to conduct the test on crystal-clear extracts. These are some-

¹¹¹ Warburg and Christian Biochem Z , 257, 492 (1933).

¹¹² Hodson and Norris J. Biol Chem., 131, 621 (1939); Scott, Hill, Norris, and Henser-J. Biol Chem., 165, 65 (1946).

¹²¹ Kuhn, Wagner-Jauregg, and Kaltschmitt Ber., 67, 1452 (1934).

times difficult to obtain when starchy samples are tested. Digestion with "Mylase" or "Taka-Diastase" is often helpful

Modified Microbiological Method of Snell and Strong: 125 Principle. Riboflavin is determined by measurement of the growth stimulation of Lactobacillus case. The acid produced by the microorganism is determined by titration with sodium hydroxide

Procedure: Preparation of Sample. Conduct all operations in dim light Weigh out a sample containing approximately 200 µ2. riboflavin and heat in an autoclave with 400 ml. 0.1 N hydrochloric acid for 30 minutes at 15 pounds pressure. Cool and adjust to pl. 14.5 with 1 N sodium hydroxide. Dilute to 1000 ml. and filter through a fine filter paper which does not adsorb iboflavin. Adjust a 100-ml. aliquot of the clear filtrate to pl. 16.8 with 1 N sodium hydroxide solution and dilute to 200 ml. Filter again if the solution is not clear,

Prepare the following solutions:

Yeast Extract Solution. Heat a suspension of 500 g. fresh, starch-free baker's yeast in 5 liters of distilled water for 2 hours in flowing steam; then autoclave for 40 minutes at 15 pounds pressure. Allow the suspension to settle, filter, and evaporate the filtrate to 125 ml, under reduced pressure below 50° C.

Yeast Supplement Solution. Add 125 ml. of an aqueous solution containing 38 g. lead subacetate to 125 ml. of yeast extract solution. Filter and adjust the filtrate to pl1 10 with ammonia. Filter and adjust the pl1 to 6.5 with glacial acetic acid. Treat the solution with hydrogen sulfide, filter off the lead sulfide, and dilute the filtrate to 250 ml. with water. Preserve

Reference Standard Solution. Dissolve 50 mg. U.S.P. ribofiavin reference standard in 500 ml. distilled water containing 1 ml. giacial acetic acid. Preserve this stock solution under toluene in an amber bottle in the refrigerator. For each set of assays, dilute 1 ml. of this solution to 1000 ml. with distilled water. This solution contains 0.1 pg. riboflavin per ml.

Basal Medium Stock Solution. Dissolve 15 g. anhydrous dextrose in a mixture of 50 ml. photolyzed peptone solution, 50 ml. cystine solution, 5 ml. yeast supplement solution, 2.5 ml. salt solution A, and 2.5 ml. salt solution B. Adjust to pH 6.8 with 1 N sodium hydroxide and dilute to 250 ml.

Preporation of the Stock Culture. Drivte 10 ml. yeast extract solution to 100 ml. with distilled water, and add 1 g. anhydrous destrose and 1.5 g. agar. Heat on a steam bath to dissolve the agar. Add approximately 10 ml. portions of the hot solution to test tubes, plug with nonabsorbent cotton and sterilize in an autoclave at 15 pounds pressure for 20 minutes. Allow to cool in an upright position. Prepare several stab cultures of Lactobacillus casei (American Type Culture Collection No. 7469) and incubate for 16 to 24 hours at any constant temperature between 30° and 37° C. Finally store in a refrigerator. Prepare a fresh stab every week, and do not use for inocultum if more than two weeks old.

Preparation of the Inoculum. To each of two tubes containing 5 ml. of the basal medium stock solution, add 5 ml. distilled water containing 1 µg. riboflavin. Autoclave at 15 pounds pressure for 20 minutes and cool, 70 one of the tubes make a transfer of cells from the stock culture of Lactobacillus casel and incubate for 16 to 24 hours at any constant temperature between 30° and 37°. Transfer 1 drop of this culture to the second tube and incubate again for 16 to 24 hours. Centrifuge the culture under aseptic conditions and decant the supernatant liquid. Suspend the cells in 10 ml. sterile isotonic solution of sodium chioride.

Assay. In duplicate tubes, 16 by 150 mm. in size, place respectively 0.5, 1.0, 1.5, and 2.0 ml. of the extract of the test material. To each add 5 ml. of the basal medium stock solution and sufficient distilled water to bring the volume in each tube to 10 ml. Prepare a similar set of duplicate tubes containing respectively 0.00, 0.05, 0.10, 0.15, 0.20, 0.30, and 0.50 µ£. standard riboflavin. Mix the solutions thoroughly, plug the tubes with nonabsorbent cotton, and autoclave at 15 pounds pressure for 20 minutes. Cool aseptically, add 1 drop of inoculum to each tube, and incubate for 72 hours at any constant temperature between 30° and 30° C. Keep all the tubers in darkness or semidarkness during their preparation and incubation, and protect against contamination by foreign microorganisms. Transfer the contents of each tube to a small Erlenmeyer flask, using a fixed volume of distilled water for rinsing. Titrate with 0.1 N sodium hydroxide using bromorhymol blue as the Indicator.

CALCUATION On ordinary graph paper, plot the average titrations in ml. of 0.1 N sodium hydrovide aguinst ag of riboflavin in the sense of standard tubes. From this standard curve, estimate the riboflavin content of each ml of the test solution in each duplicate set of tubes. Calculate the riboflavin content of the test material from the average values obtained from not less than three sets of these tubes which do not vary by more than ±10 per cent from the average.

Comment. The microbiological method for the determination of riboflavin employing Lactobacillus casei is highly sensitive and shows a good degree of specificity for the vitamin If desired, turbidity in place of acidimetric titration may be used to measure response. For measurement of turbidity follow the directions given in the Microbiological As.as for Vitamin B11, p 1210, beginning with the fifth paragraph under the heading Procedure and substituting riboflavin for cyanocobalamin For this procedure it may be necessary to reduce the level of riboflavin used in the standard and sample tubes

Fluorometric Method of Arnold 184 Principle The fluorometric procedure for the determination of riboflavin depends upon the extraction of the vitamin with dilute acid filtration treatment of the filtrate with permanganate and hydrogen peroxide to destroy interfering pigments and measurement of the fluorescence. The vitamin content of the extract is evaluated by means of an internal standard

Procedure Suspend a finely ground sample containing approximately 5 pg riboflavin in 75 ml 0 1 N sulfuric acid Heat in an autoclave for 15 minutes 21 15 pounds pressure, or in a boiling water bath for 45 minutes with intermit tent shaking Cool the suspension and adjust the pil to 4 3 with 2 5 M sodium acetate Dilute to 100 ml , shake well, and filter through Whatman No 1 or No 40 filter paper Discard the first 15 ml filtrate Treat 60 ml of the filtrate with 2 ml 4 per cent potassium permanganate solution After 3 minutes, dis charge the permanganate color with freshly prepared 3 per cent hydrogen peroxide solution Break the froth with a few drops of acetone, and dilute to 65 ml with distilled water Mixand filter Pipet out two 15 ml portions of the filtrate To one, add 1 ml distilled water and measure the fluorescence (A) (see p 1163) To the other, add i mi riboflavin standard solution containing 1 µg of the vitamin, and measure the fluorescence (B) Obtain the blank (C) after adding exactly 20 mg of sodium hydrosulfitein to A or B

CALCULATION Obtain the riboflavin content of the sample using the formula,

$$\frac{A-C}{B-A} \times \frac{1}{15} \times \frac{65}{60} \times \frac{100}{G} = \mu {\rm g}$$
rīboflavin per g_of sample

where G is the weight of the sami le taken and A, B and C are as indicated above

Comment. The fluorometric method may be employed for the determination of the riboflavin content of yeast, white flour, bread, milk powder, and similar products However, for samples which yield highly pigmented extracts, or for materials containing less than I µg of riboflavin per g of solids the microbiological procedure is preferred

Determination of Riboflavin in Urine (Flurorometric Method of Najjar) Principle The risoflavin is extracted with acetic acid pyridine-butanol mixture after interfering urmary pigments are oxidized with permanganate. The concentration of mboffax in in the extract is measured fluorometrically

Procedure Collect a 24 hour urine sample in a brown glass bottle contain ing 20 ml of 10 per cent sulfuric acid Since riboflavin is destroyed by hightconduct all of the following operations in semidarkness Measure the volume

¹⁸ Arnold Cereal Chem 22, 455 (1945)

¹²⁷ Solium bydrosulfite (highest purity) al ould be stored so as to avoid undue exposure to

in \ajjar J Biol Chem 141 355 (1941)

Transfer an aliquot containing approximately 5 µg riboflavin to a 100 ml centrifuge cup, and add sufficient water to make the volume 10 ml Add 2 ml glacial acetic acid and 4 ml coloriess pyridine and mix For each ml of urine, add 2 drops 4 per cent KVinO₄ After exactly 1 mlnute, decolorize the excess permanganate with a few drops of freshly prepared 3 per cent hydrogen peroxide solution Add 10 g anhydrous sodium sulfate, then 20 ml n butanol Shake videorously for 5 minutes Centrifuge

Determine the fluorescence of the supernatant solution, that of a stand ard containing 5 µg riboflavin, and of a water blank, the latter two likewise carried though the entre procedure.

For these measurements, a Pfaltz and Bauer fluorophotometer may be employed. The source of light should be a mercury vapor lamp equipped with a dark blue filter (Jena, No. BG12) and a yellow filter (Jena No. GG3). An orange filter (Jena, No. OG1) should be placed between the fluorescent solution and the photocell. Set the diaphragm so that the galvanometer reads 50 when the extract of the 5 mg standard is in the instrument.

At the concentrations recommended in this procedure, the fluorescence of a riboflavin solution is directly proportional to its concentration. With most instruments, the galvanometer deflection is proportional to the fluor escence. Establish the linearity of the instrument employed by measuring the fluorescence of standard aqueous solutions of riboflavin as follows.

Dissolve 50 0 mg riboflavin in 1000 ml of hot water in the dark. Cool Prepare aqueous dilutions containing 0.00, 0.05, 0.10, 0.15, 0.20, 0.25, 0.30, and 0.40 µg riboflavin per ml Set the instrument at 100 with the most concentrated standard and measure the fluorescence of the other solutions. Plot the galvanometer readings against concentration of the witamin on ordinary graph paper. Check the settling of the instrument before each reading.

CALCULATION If the relation is linear, calculate the riboflavin content of the 24-hour sample by means of the following formula

$$\frac{U-C}{50-C} \times 5 \times \frac{V}{A} = \mu g$$
 riboflavin per 24-hour sample

where U is the galvanometer reading of the unknown C is that of the reagent blank V is the volume of the 24-hour sample in ml and A that of the aliquot taken for test

If the curve obtained above is not linear standard solutions containing 0, 2.4.6 as $y_{\rm H}$ to loady must be carried through the entire procedure and the concentration of nboff avia in the unknown determined from a standard curve. This value multiplied by $\frac{V}{4}$ gives the amount of riboff axio in the 24-hour urine sample

Comment. The fluorometric method for the determination of riboflavia in urine, as described above, measures in addition uroflavia, a fluorescent metabolite of the vitamia For diagnostic purposes however, the total fluorescence following the ingestion of a test dose of the vitamia is a good index of the nutritional state with respect to riboflavia.

BIOLOGICAL ASSAY OF RIBOFLAVIN

The essential feature in the biological assay for riboflavin is to select a basal ration which includes all of the required factors of the vitamin B complex with the sole exception of riboflavin. In the past the practice has been to supplement a basal ration otherwise free from all sources of the vitamin B complex with alcoholic extracts of wheat germ or of rice polish

ings (so-called tiki-tiki). However, such diets were undoubtedly deficient in more respects than riboflavin alone. Since crystalline preparations of each of the known B vitamins are now available, it is possible to supplement an otherwise adequate diet of purified components with all of the factors of the B complex except the one under assay.

There is at present no officially recognized procedure for the biological assay of riboflavin Procedures employing both the rat and the chick have heen described The curative rat assay is more widely employed in the evaluation of foods or pharmaceutical products for human use. The customary procedure is to place rats on the riboflavin-free basal diet at weaning, making observations of weight changes at semiweekly intervals Cossation of growth is one of the earliest symptoms of rihoflavin depletion, and if the deprivation is complete, it is usually followed by decline in weight and death without the appearance of characteristic symptomatology If the deficiency is prolonged (as may occur if the hasal diet is not completely devoid of rihoflavin) the symptoms of ariboflavinosis appear These include bilaterally symmetrical patches of baldness around the nose, neck, back, and abdomen, and corneal vascularization, and if the deficiency is particularly prolonged, entaraets are observed. It is interesting to point out that nboflavin-deficient rats are particularly susceptible to pediculosis The occurrence of alopecia is noted in deficiencies of other factors of the B complex-eg, hiotin Whether this phenomenon has a common origin in deficiencies of various memhers of this group of vitamins is not known.

After three to four weeks on the riboflavin-free diet, the weight of the rat reaches a plateau or decline and at this point the material to be assayed is fed as a supplement to the basal ration Comparable groups of animals receive graded dosages of crystalline riboflavin at levels which result in a graded dose-response curve. For example, doses of 1, 2, 4, 8, and 16 gg of crystalline riboflavin may he fed daily. Together with a resumption of growth, restoration of the body hair occurs over the denuded areas Potency may be estimated by comaprison of the log dose vs. growth response data for the test material with those for the standard (dealculation in vitamin A growth assay, Appendix VI)

NIACIN (NICOTINIC ACID)

The most common acute deficiency disease in this country is pellagra. Before the discovery of its relationship to macin, this disease was responsible for thousands of deaths annually in the South Pellagra occurs in endemic and epidemic form in many other countries including the Soviet Union, Egypt, Italy, Spain, and the Balkan countries Epidemic pellagra follows any serious interruption of food supply or intake, especially after war, famine, disease, or economic depression.

Pellagra has been known for many centuries and was described as early as 1735 by Casal of Spain That the disease was of dietary origin was not recognized until 1912 when Goldberger, Waring, and Willets¹³ demonstrated that pellagra could be prevented by an adequate diet. At about

⁹ Goldberger, Waring, and Willets U.S Pub Health Repts . 30, 3117 (1915)

the same time, nicotinic acid, a compound synthesized as early as 1867, ¹³⁰ was isolated by Funk while attempting to purify the antiberiberi vitamin. He failed to recognize the relationship between nicotinic acid and pellagra but noted its beneficial effects when fed along with "vitamine B." Deficiency syndromes analogous to human pellagra were demonstrated in dogs¹³² in 1917 and in albino rats¹³³ in 1926. It was soon recognized that those foods which were effective for the cure of caninc blacktongne were also effective for the cure of pellagra. ¹³⁴ Following the discovery in 1937 of the importance of the amide of nicotinic acid (niacinamide) in the nutrition of certain unicellular organisms, ¹³⁵ the therapeutic effects of nicotinic acid in the treatment of blacktongue¹³⁶ and of niacinamide in pellagra. ¹³⁷ were demonstrated.

Physiological and Clinical Aspects of Niacin. In piacin deficiency the demonstrable tissue lesions are preceded by functional disturbances which are characteristic of deficiencies of a number of the B ritaming These include weakness, anorexia, indigestion, diarrhea, and mental and emotional disturbances. Though pellagra is generally complicated by deficiencies of other B vitamins, certain lesions of the skin, the directive tract, and the nervous system (the "three D's"-dermatitis, diarrhea dementia) have been attributed specifically to deficiency of niacin. The skin lesion is an erythema, generally bilaterally symmetrical, which affects the back of the bands, the knees, elbows, dorsum of the feet, and ankles. Exposure to the sun increases the severity of the lesions. In chronic pellagra the affected areas become permanently pigmented and either atrophied or roughened and thickened. Epithelial lesions are also noted in the digestive tract, particularly on the tongue (glossitis) and in the rectum. The organic nerve lesions of pellagra include myelin degeneration of the spinal column fibers and degeneration of the axis cylinders of pyramidal cells of the cortex. Mild mental disturbances are common in nellagra, and acute delirium and dementia occur in severe cases.

Though anemia is not generally recognized as a symptom of pellagra, it is frequently accompanied by a macrocytic anemia. Since this condition responds to pteroyfglutamic (folic) acid, its occurrence in pellagra may be due to the multiple nature of the clinically observed condition. Dogs with blacktongue also show anemia which responds to the administration of naicin. 1.18

Dramatic cures of pellagra occur upon the administration of niacinamide. Therapeutic doses of 50 to 500 mg, of niacinamide per day cause the disappearance of glossitis in 24 hours and of lesions of the tonnue and

¹³⁰ Huber: Liebig's Ann. Chem., 141, 271 (1867).

¹³¹ Funk: J. Physiol., 46, 173 (1913); Brit. Med. J., 1, 814 (1913). Drummond and Funk: Biochem. J., 8, 594 (1914).

¹³¹ Chittenden and Underhill: Am. J. Physiol., 44, 13 (1917).

Goldberger and Lillie: U.S. Pub. Health Repts., 41, 1025 (1926).
 Aykroyd and Roscoe: Biochem. J., 23, 483 (1929).

¹⁴⁵ Knight: Biochem. J., 31, 731 (1937). Muller: J. Biol. Chem., 120, 219 (1937).

Tivelijem, Madden, Strong, and Woolley: J. Am. Chem. Soc., 59, 1767 (1937); J. Biol. Chem., 123, 137 (1938).

I'll Pouts, Helmer, Lepkovsky, and Jukes: Proc. Soc. Exp. Biol. Med., 37, 405 (1937).
 Handler and Featherston: J. Biol. Chem., 151, 395 (1943).

lips in three to five days Since uncomplicated niacin deficiency is comparatively rare, other members of the vitamin B complex are usually also administered in treating the disease

Niacinamide is an essential constituent of diphosphopyridine and triphosphopyridine nucleotides (DPN and TPN, also known as coenzymes I and II, or codehydrogenases I and II), which participate in a large number of physiologically important oxidation reactions. These coenzymes occur in practically all cells and have been synthesized from niacin and niacinamide in utro by nucleated cells. The structures of coenzyme I and of its reduction product, reduced coenzyme I, are shown below:

The structure of reduced coenzyme I is based upon the evidence presented by Pullman et al. 1300 that reduction involves saturation of the carbon atom para to the introgen in the pyridine ring, rather than ortho to it, as had been previously thought Note furthermore that reduction involves the addition of only one hydrogen atom to a molecule of coenzyme; the second hydrogen furnishes an electron to the quaternary N and becomes a hydrogen ion. Hence the stochhometry of the reaction is as follows.

TPN differs from DPN in that it contains an extra phosphoric acid group probably linked to the 2' position of the ribose mosety of adenosine These coenzymes, in the presence of specific proteins, catalyze physiological oxidation reactions Among the 35 different known reactions in which DPN participates are the oxidation of alcohol to acetaldehyde, of glucose to gluconic acid, of mahe acid to oxalacetic acid, of lactic acid to pyruvic acid, and of gly cerophosphate to phosphoglyceraldehyde Among the enzyme reactions catalyzed by TPN are the conversion of Robison's ester (glucose-6-monophosphate) to phosphochyonic acid and of glutamic

¹³⁴⁴ Pullman, San Pietro and Colowick J Biol Chem , 206, 129 (1954)

acid to iminoglutamic acid. These enzymatic reactions are all reversible. and the reduced coenzymes may combine with their anoenzymes to catalyze physiological reductions. In practice there is a dynamic equilibrium which depends on the relative concentrations of the reduced and ovidized forms of the substrates and coenzymes and on other conditions. In vitro experiments indicate that reduced coenzymes I and II may be reoxidized by flavoprotein enzymes, and that coenzyme II may be converted to I by a phosphatase 139 Roth coenzymes are inactivated by enzymetic destruction when cells of brain, liver, kidney, or muscle are runtured. In contrast to the riboflavin coenzymes, which are present in flavorroteins in simple numerical proportion to their appearzymes, the piacinamide coenzymes are found in great excess and are bound only loosely to the protein moieties. The macinamide prosthetic groups have been designated mobile ecentymes. DPN can be hydrolyzed by dilute alkali the cleavage occurring between the niacinamide and ribose mojetics. Similar splitting can be induced by enzymes (designated DPN nucleosidases) found in brain tissue140 and in Neurospora 141

That dietary niacin is important in the maintenance of physiological enzyme systems is indicated by the positive correlation between the intake of the vitamin and the concentration of the coenzymes in the muscles 142 Niacin is essential to the physiology of animals plants and microorganisms. All plants and some microorganisms synthesize the vitamin. Others-e.g., C. diphtheriae, lactic acid bacteria, and B. dusenteriae-require an external source, and still others-e.g., B. influenzaerequire an external source of the coenzymes. Though the mold Neurospora does not ordinarily require niacin, a mutaut strain produced by irradiation requires an external supply of the vitamin. Whereas niacin is essential to all animals, certain species, e.g., the rat and the horse, do not require dietary sources because their intestinal flora synthesize enough to meet metabolic needs. A dietary supply is required by the cotton rat, dog, pig, rabbit, chick, monkey, and man. However, even in these species, significant contributions are made by intestinal microorganisms. Urinary excretion studies143 show that human subjects on a constant diet excrete lower concentrations of N¹-methylnieotinamide (the principal metabolite of niacin and niacinamide found in human urine) after dosage with sulfacuanidine and succinvlsulfathiazole which destroy intestinal bacteria. Microorganisms have been isolated from the human cecum144 which synthesize niacin in vitro.

The amount of miacin required by animals from external sources depends upon the nature of the diet, particularly with regard to protein and carbohydrate. Endemie pellagra oecurs almost exclusively in localities where corn is employed as a staple cereal. Populations which consume

144 Benesch: Lancet, 1, 718 (1945).

¹⁴⁹ Euler and Adler: Z. physiol. Chem., 252, 41 (1938).

¹⁴⁰ Gore, Ibbott, and McIlwain: Biochem. J., 47, 120 (1950).

Kaplan, Colowick, and Nason: J. Biol. Chem., 191, 473 (1951).
 Anderson, Teply, and Elvehjem: Arch. Biochem., 3, 357 (1944).

¹⁴⁹ Ellinger, Coulson, and Benesch: Nature, 154, 270 (1944); Ellinger, Benesch, and Kay: Laneet. 1, 432 (1945).

large quantities of rice rarely show pellugra even though their dictary intake of macin is no more than 5 mg per day, whereas mane-eaters develop pellugra despite much higher intakes 145 Rats which ordinarily synthesize their own macin show a typical macin deficiency syndrome when fed acorn dict 146 This may be prevented by adding either tryptophan or more protein On an otherwise adequate dict young growing pigs show little sign of macin deficiency, though the amount of the vitamin ingested is small. A decrease in the protein of the ration however, produces symptoms of macin deficiency, which may be prevented by administration of more of the vitamin.

Since macin deficiency in white rats on a corn diet is prevented by the administration of tryptophan it appears that the lack of this amino acid which is present in only low concentrations in corn is responsible for the apparent increased requirement of many species for macin. This effect may possibly be attributed to changes in the intestinal flora resulting in the loss of microorganisms which require tryptophan from which they are capable of synthesizing macin. Tryptophan serves as a precursor for the hiosynthesis of macin not only by these microorganisms but in animal tissues as well. Hence pellagra must be viewed as a dual deficiency of both tryptophan and macin. The possible existence of a pellagragenic antivitamin (see Chapter 36) in corn has received consideration.

The effect of corn on the growth of white rats depends upon the nature of the carbohy drate ingested. When the latter is sucrose considerable in hibition of growth occurs. However when sucrose is replaced by glucose and dextrin or partially replaced by lactose the inhibition of growth does not occur. 14 Further evidence that the increased requirement for macin is associated with the nature of dietary carboby drate and with a dimmished intake of tryptophan is provided by observations on white rats subsisting on a diet containing wheat gluten and gelatin and deficient only in macin and tryptophan. With sucrose as the source of carbohydrate very poor growth resulted. The administration of tryptophan or macin or the substitution of glucose and dextrin for sucrose restored normal growth.

The principal metaholite excreted in the urine of man rats dogs and cats following the ingestion of maein or macinamide is \(^1\) methyl meotinamide Some mootinamide but only small amounts of meotinic acid are also excreted. However rabbits and guinea pigs excrete principally meotinic acid and practically no \(^1\) methylmicotinamide \(^{10}\) Fle latter compound after alkaine hydrolysis reacts like macin with cyanogen bromide and amline and because of its physical and chemical similarity to trigonelline (the betaine of \(^1\) methylmicotinamide has the following structure.

Pellagrins were observed to exercte only small quantities of this metabolite Early tests for the diagnosis of ninein deficiency were based upon the colorimetric determination of urinary "trigonelline" after the ingestion of a test dose of the vitamin. It was subsequently discovered in that administration of piacin or piacinamide was followed by exerction of a metabolite which was readily converted to a highly fluorescent substance by alkalinization and subsequent extraction with butanol. The urine of nincin-deficient subjects contained only small amounts of this compound. even after dosage with the vitamin. The metabolite was identified as Ni-methylnicotinamide152 and found to be identical with the compound measured colorimetrically after alkaline hydrolysis which had been erroneously designated "trigonelline." Humans in a normal state of nutrition excrete approximately 20 per cent of their intake of niacinamide in the form of the N1-methyl compound on a molar basis and approximately half of this amount when ingesting macin. 154 The conversion of the latter to N1-methylnicotinamide requires both an amidation and a methylation. whereas niacinamide requires only the latter step. In ritro studies154 show that rat liver (but not kidney or muscle) is capable of converting niacinamide, but not niacin, to N1-methylnicotinamide, the degree of conversion being increased by the addition of methionine.

The possibility that macin or its amide acts physiologically in the form of N-metbylnicotinamide has been investigated with conflicting results. Some observers have found that the compound is capable of curing blacktongue, 155 but this has been disputed by others. 156 In man, N'-methylnicotinamide has been found to improve the dermatitis and glossitis of pellagra but simultaneously to aggravate the psychomotor symptoms. 157 The latter however, could be alleviated by the administration of thiamine and riboflavin. In view of these findings, the metabolic role of N1-methylnicotinamide remains to be established.

Certain derivatives of niacin and niacinamide are biologically active in higher animals when administered orally. These include various esters, and the ureide and diethylamide of niacin. These compounds are probably

¹¹¹ Najjar and Wood: Proc. Soc. Exptl. Biol. Med., 44, 386 (1940); Najjar: Bull, Johns Hopkins Hosp., 74, 392 (1944).

¹³² Huff and Perlaweig: Science, 97, 538 (1943); J. Biol. Chem., 150, 395, 483 (1943),

Hochberg, Melnick, and Oser: J. Biol. Chem., 138, 265 (1945).
 Perlzweig, Bernheim, and Bernheim: J. Biol. Chem., 150, 40 (1913); Ellinger: Biochem

J., 42, 175 (1948).

Najjar, Ilall, and Deal: Bull, Johns Hopkins Hosp., 76, 83 (1945).
 Tepley, Krehl, and Elvehjem: Proc. Soc. Exptl. Biol. Med., 58, 169 (1945).

¹¹⁷ Vance: Bull. Johns Hopkins Hosp., 77, 393 (1915).

large quantities of rice rarely show pellagra even though their dietary intake of maein is no more than 5 mg per day, whereas maize-eaters develop pellagra despite much higher intakes 148 Rats which ordinarily synthesize their own maein, show a typical maein deficiency syndrome when fed a corndict 148 This may be prevented by addingetther tryptophas or more protein. On an otherwise adequate diet, young growing pigs show little sign of maein deficiency, though the amount of the vitamin nigested is small. A decrease in the protein of the ration, however, produces symptoms of maein deficiency, which may be prevented by administration of more of the vitamin.

Since macin deficiency in white rats on a corn diet is prevented by the administration of tryptophan, it appears that the lack of this amno acid, which is prevent in only low concentrations in corn, is responsible for the apparent increased requirement of many species for macin. This effect may possibly be attributed to changes in the intestinal flora resulting in the loss of microorganisms which require tryptophan from which they are capable of synthesizing nacin. Tryptophan serves as a precursor for the biosynthesis of macin not only by these microorganisms but in animal tissues as well. Hence pellagra must be viewed as a dual deficiency of both tryptophan and macin. The possible existence of a pellagragenic antivitation (see Chapter 36) in corn has received consideration.

The effect of corn on the growth of white rats depends upon the nature of the carbohydrate ingested. When the latter is sucrose, considerable in libition of growth occurs. However when sucrose is replaced by glucose and dextrin, or partially replaced by lactose the inhibition of growth does not occur. In Turther evidence, that the increased requirement for macin is associated with the nature of dictary carbohydrate and with a dimmisshed intake of tryptophan is provided by observations on white rats subsisting on a dict containing wheat gluten and gelatin and deficient only in macin and tryptophan. With sucrose as the source of carbohydrate, very poor growth resulted. The administration of tryptophan or macin or the substitution of glucose, and dextrin for sucrose, restored normal growth.

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Najjar, Hall, and Deal Bull Johns Hopkins Hosp., 76, 83 (1945)
 Tepley, Krehl, and Hychyem Proc. Soc Exptl Biol. Med., 58, 169 (1945).

¹¹⁷ Vance Bull, Johns Hopkins Hosp , 77, 393 (1915)

hydrolyzed in the gastrointestinal tract to macin and macinamide Trigonelline, the betaine of N-methylmacin, is biologically mactive It occurs naturally in a number of foods, especially legumes, coffee, and tobacco A synthetic analog of macin, pyridine-3-sulfonic acid, has been found to inhibit the growth of microorganisms requiring macin or its amide 158 3-Acetylpyridine does not inhibit microbial growth but in duces macin deficiency in mice, a species which normally does not require a dictary supply of macin 155 This compound, becau-e of its chemical similarity to macin, probably acts by competition with the vitamin in plis stological enzyme systems (see Chapter 36)

Wheat bran and other natural materials contain a compound which upon heating with strong acid or alkali, heliaves like macin toward Lactobacillus arabinosus 100 This compound is extracted from natural ma tenals by heating with water or dilute acid, but must be subsequently hydrolyzed with strong acid or with alkali before it can stimulate the microorganism Concentrates of the unhydrolyzed material have been prepared and found to be mactive when fed to chicks After hydroly is the compound meets the macin requirements of both the chick and the dog Nikethamide ("Coramine"), macin-diethylamide, exhibits similar behavior 141 in that it is mactive for L arabinosus but is active after heating with alkali Nikethamide is biologically active in higher animals

I arge do-es of macin dilate the superficial blood tessels. The administration of 50 mg or more orally or of 10 mg intravenously to humans not deficient in the vitamin produces flushing, itching, and burning of the skin especially in areas subjected to pressure Individuals vary greatly in their sensitivity to this reaction. The effect is observed only with the free acid and not with macinamide, so that the latter is preferred for thera peutic purposes The flushing reaction may be prevented by simultaneous administration of gly cine which provides amino groups for conversion of

the free vitamin to its amide

Large doses of macin or macinamide are toxic to the rat because of the depletion of the methyl donor, methionine 14 Nacinamide is more touc than the free acid since more \1-methylmicotinamide is excreted when the amide is fed The toxicity is seen only in animals which excrete Vi-methylmicotinamide after do-age with the vitamin Large do-es of macmamide are not toxic to young rabbits and guinea pigs, which do not excrete the methylated derivative The toxicity of large do-es of the vita min in rats may he prevented by feeding methionine, or choline plus homoeystine (see p 1030)

For further discussion of the clinical aspects of macin, see the American Medical Association syllabus p 1291

Storage and Synthesis of Niacin. No significant amount of niacin of its amide is stored passively in animal to use Higher concentrations are

Melliam Brd J Expl Path 21 13F (1940)
 Woodley J Biol Chem 157 4aa (191a)

nomey 3 Biol Chem 157 Aug (191a)

118 Andrews Boyd and Gortner Ind. Eng Chem Anal Pd. 14 653 (1942) Krehl and

119 Story J Biol Chem 154 (1944)

11 Tesh) and Elvel jetta Proc Sur Payal Biol Med. 55 °2 (1944)

11 Handler J Pol. Ch. 155 (1945)

¹¹¹ Handler J Bud Chem 154, 203 (1944)

found in the liver, muscle, and kidney for the performance of metabolic functions. A decrease in the macin intake of man is followed by a diminished concentration of the macin coerzymes in the striated muscles, but has little effect on the spenzyme content of the crythrocytes.

Biosynthesis of macin by plants and microorganisms, the importance of the contribution by intestinal flora in the nutrition of animals, and the effect of other dietary factors have been discussed above

A number of practical syntheses for mrem and macmamde are available for commercial use. The simplest of these include the nitric acid oxidation of mootine obtained from tobasec or the oxidation of β procline or quinoline with potassium permanganate or in the vapor phase with air and metallic oxide catalysts. Nincinimide may be prepared from the free acid by heating with ammonia, from esters by treatment with alcoholic ammonia, or from 3 evanopridine by acid or alkaline bydrolysis.

in the presence of hydrogen perovide and subsequent treatment with a

quaternary ammonium resin is Distribution of Niaclin. Is In natural materials, macin occurs predominantly as the amide, the form in which it is active biologically in enzyme systems. Good sources of the vitamin are liver, adrenal glands, fish, meats, whole wheat or rye, and enriched flour. The vitamin occurs in relatively high concentration as coenzyme I in yeast, red blood cells, and beart muscle. Fruits are poor sources of the vitamin muts are somewhat incher, and vegetables are variable potatoes and legumes being generally higher than leafy varieties. Con's milk may contain from about I mg to as much as 5 mg per quark, whereas human milk contains rela

tively less Ten, coffee and beer are good sources of macin Chemistry of Niacin Niacin (pyridine-3-carbox) he neid) is a white crystalline solid which melts at 236° C and may be sublimed without destruction The vitamin is stable to boiling in neutral and or alkaline solution. The vitamin has an absorption maximum in the ultraviolet region at 385 mu.

$$\bigcap_{N}^{O} \stackrel{O}{=} C-NH$$

Niacin (Nicotinic Acid)

Niacinamida

Nincinamide crystallizes in light needles from benzene and melts at 129° C. It may be distilled at 150 to 160° at 5 × 10 4 mm. Nincinamide is more soluble in organic solvents than the free acid and inhibe the latter, may be extracted from water with ether. The amide has a characteristic absorption curve in the ultraviolet region with maxima at 210, 220, and 260 mm. Niacinamide is readily hydrolyzed to free macin by heating with acid or all shi

Niacin and miacinamide react with cyanogen bromide in the presence of

¹⁶³ Galat J Am Chem Soc 70 3945 (1948)

¹⁴⁴ See Appendix III

a primary or secondary amine to produce a compound having a yellowgreen color Colors are also formed with trigonelline, nicotinuric acid, and nicotine, as well as certain other pyridine compounds. This reaction is employed in the chemical determination of macin. Since the amide produces only half the color of the free acid, it must first be hydrolyzed for

the colorimetric te-t Coenzymes I and II (DPN and TPN) are colorless, water-coluble com pounds in coluble in organic colvents Coenzyme II, however, is soluble in methanol-hydrochloric acid mixtures Both coenzymes are relatively stable in acid solution but are readily decomposed by alkali Coenzyme l has an absorption maximum at 260 mg. It is nonfluorescent, optically active, [a] 540 m $\mu = -70^{\circ}$, and is not precipitated by lead acetate, but does produce an in-oluble cuprous salt. Reduced coenzyme I has an absorption maximum at 340 mg and fluoresces upon ultraviolet irradiation This compound is stable in alkali but is destroyed by acid in which it reverts to the oxidized form which is non-fluorescent. Coenzy me II has an absorption maximum at 360 mg. Its optical rotation [a] 546 mg = -29.4 Its cuprous salt is soluble but the coenzyme is precipitated by lead acetate

Determination of Niacin. Accurate chemical and microbiological method, for the determination of macin in natural materials are available able The chemical procedures involve reaction of the vitamin with cyanogen bromide in the presence of an amine Extracts are prepared for chemical or microbiological assay by acid or alkaline hydrolysis, hoth to facilitate extraction and to hydrolyze macinamide and a macin precursor found in natural materials. In the chemical procedure, hydrolysis with acid results in the formation of furtural from the degradation of pentowhich interferes with colorimetric reaction. This compound is esparated from the macin by adsorption of the vitamin on a hydrated aluminum silicate which does not adsorb the furfural. The macin is subsequently eluted from the adsorbate by washing with alkali

Because macin and macinamide have somewhat different physiological effects, it is sometimes desirable to determine their relative concentrations in mixtures This may be accomplished chemically by time-reaction measurements conducted on the hydrolyzed and unhydrolyzed extracts it The assay is based on the differences in the rate of formation of the yellow-

green pigment with cyanogen bromide and aniline

The microbiological determination of macin is based upon the growth stimulation of Lactobacillus arabinosus. This organism responds equally to macin and its amide These compounds, however, may be differentiated microbiologically by destruction of the amide with bromine and potassium hydroxide (Hoffman degradation) converting it to β -amino-pyridine which is microbiologically inert is Differential microbiological assays may also be conducted to distinguish between the several metabolites of the vitamin, including macin, its amide micotinume acid, and Numethal nicotinamide Two microorganisms are employed, Lactobacillus arabino-

¹⁴ Lamb Ind Eng Chem Anal Pd 15 252 (1943) Melnick at l Over Jul., 15 34 10 Atkin Schultz Williams and Frey J Am Chem Sor 65 992 (1947)

sus, which utilizes niacin, nineinamide, and nicotinuric acid, and Leuconsider mesenteroides, for which only piacin is active. 167 Hydrolysis with 0.6 N sulfure acid at 15 pounds pressure for one hour is also employed to convert the amide completely to free ninein without affecting nicotinurie acid. The microbiological assays are conducted in conjunction with the fluorometric test which measures only N1-methylnicotinamide Differential assays such as those described above show that the major part (about 90 per cent) of the miscin metabolites excreted after the ingestion of the vitamin appears as N1-methylnicotinamide. Small amounts of practice and practice also occur in urme and somewhat larger amounts of the latter after the administration of large doses of the free acid. Human urine contains little if any nicotinuric acid except after large doses of free nicotinic acid

The biochemical diagnosis of ninein deficiency is based upon the measprement of prinary excretion of N's-methylnicotinamide 168 A satisfactory clinical test involves measurement of the 6-hour urinary excretion of N1-methylnicotinamide following the oral administration of 300 mg nigeinamide 189

Earlier clinical tests for the diagnosis of pellagra, based upon measurement of urmary "porphyrins," are not reliable since the color reaction has been found to be due to prorosein rather than to porphyrin and denends on the presence of other compounds in the urine

Modified Colorimetric Method of the Research Corporation:170 Principle. The method involves strong acid hydrolysis to convert derivatives to nicoting acid. adsorption of the vitamin on Lloyd's reagent, clarification of the cluste with lead hydroxide, and reaction of picotinic acid with evanogen bromide and amiline to produce a vellow pigment which is measured photometrically.

Procedure: Preparation of the Sample, Hydrolyze a sample of tissue or food containing 100-500 pg. of niacln by heating In a boiling water bath with 75 ml. 4 N hydrochloric acid for 30-40 minutes. Gool and dilute to 100 ml. Transfer 25 ml, of the suspension to a narrow (18-mm, diameter) centrifuce tube calibrated at the 26.5-ml. mark. Adjust the pH to 0.5-1.0 with 18 N sodium hydroxide using 0.1 per cent methyl violet as an outside indicator. 171 Cool, add 2.5 g. Lloyd's reagent, 172 and shake I minute. Centrifuse and discard the supernatant liquid. Wash the residue twice with 10 mi. 0.2 N suifuric acid, each time centrifuging and discarding the supernatant, Add 0.5 N sodium hydroxide to the 26.5-ml, mark and break up the precipitate with a stirring rod. Immerse the tube in a boiling water bath for 5 minutes, stirring the contents occasionally. Cool and centrifuge. Transfer the supernatant to another tube, add 1.6 g. finely divided lead nitrate, and shake vicorously for I minute. If the solution is still alkaline, add more lead nitrate until acid.

Johnson J Biol Chem., 159, 277 (1945)
 Huff and Perlzweig J Biol Chem., 150, 395 (1943)
 Goldsmith Arch Internal Med. 73, 416 (1944)

¹⁷⁰ Melnick Cereal Chem , 19, 553 (1942)

¹⁷¹ Use one drop of sample to one drop of indicator on a spot plate. Match with the blue color given by one drop of 0 2 N sulfuric acid

¹⁷¹ A by drated aluminum scheate obtained from I is Lally and Co., Indianapolis 1nd When suspended in water, 25 g occupies a volume of 15 ml

Centrifuge and pour the clear supernatant into another tube. Neutralize with tertiary potassium phosphate and 20 per cent phosphoric acid.

Colorimetric measurements are made in a photoelectric photometer. Two center settings are necessary, one for evaluating the residual color in the test solution, the other for the color developed in the chemical reaction. Set the photometer at 100 per cent transmittance with 3 ml. water and 7 ml. alcoholic huffer. 11 With this setting read a solution of 3 ml. of the test tract plus 7 ml of alcoholic buffer (A). Now set the photometer at 100 per cent transmittance with the reagent hlank: 3 ml. water, plus 6 ml. cyanogen bromide solution 12 plus 1 ml. aniline solution. 13 Remove the tube and record the center setting. To 3 ml. of the test extract, add 6 ml. cyanogen bromide, and 10 mnutes later 1 ml. aniline; read the solution 5 minutes later using the center setting obtained with reagent blank (B). Similarly read a solution containing 3 ml. of test extract, 10 gd. of nicothica acid in 0.1 ml. of an alcoholic solution, 6 ml. of cyanogen bromide, and 1 ml. of aniline (C).

CALCULATION Convert galvanometer readings, G, to photometric density, P,D, as follows.

$$PD = 2 - \log G$$

Then

$$\frac{B-A}{1\,01C-B} \times 10 \times \text{dilution factor} = \mu\text{g}$$
 meeting acid per g sample

Comment. Values obtained by the colorimetric method for nicotinic acid agree well with those obtained by microbiological assay of acid or alkaline extracts. Exhaustive aqueous extraction, without preliminary hydrolysis, removes all the meotine acid from cercal products but yields lower microbiological and chemical values. A meotinic acid derivative other than meotinamide is broken down by the hydrolytic procedure (see p. 1172).

column by washing with water (see p 1145). Pass a standard solution con taining 50 µg, of N1 methylnicotinamide chloride¹⁷ in 50 ml acetate buffer through the column in the same manner as

through the column in the same manner a the diluted urine samples

Fluorometric Measurement Mix the cluate and pipet a 5 ml allquot into each of two 30 ml separatory funnels To one, the blank, add 1 ml of water followed by 16 5 ml n-buta nol To the other, the test, add 16 5 ml buta nol, then 1 ml 15 per cent sodium hydroxide Stopper the vessels immediately and shake vigorously for 3 minutes Centrifuge for 0 5 minute and discard the aqueous layer Clarify the butanol by shaking with approximately 1 a anhydrous sodium sulfate Allow the butanol extracts to stand in the dark for 15 minutes during which the fluorescence reaches a maximum, then read in a fluorometer with appropriate light filters. Use a solution of quinine suifate containing 0 2 ug

per ml in 0.1 N sulfuric acid to check the setting of the instrument CALCULATION Calculate the 24-hour unnary exception of N-methylincotinamide chloride or pressed in terms of magnamide using the formula,

$$50 \times \frac{G_u}{G_s} \times \frac{60}{M} \times 0$$
707 = $\mu \mathrm{g}$ niacinamide per 24 hr

G, and G, are the galvanometer deflections of the unknown and the stundard both corrected for their respective blanks M is the period represented hy the urinary aliquot expressed in minutes and 0.707 is the factor for conversion of N-methylmicotina mide chloride to macinimide

Comment N¹ Methylmeotmamide is the principal derivative of macinamide found in human urine Average basal excretion values approximate 25-30 per cent of

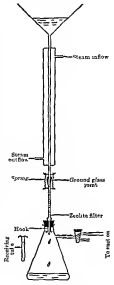


Fig 278 Apparatus for the adsorption and elution of A¹-methy luicotinamide in urine

the dietury intake with wide individual variations. The metabolite is excreted rapidly following ingestion of the vitumin approximately 20 per cent appearing in the urine within 24 hours following docage with 50-200 mg macinamide.

the material to be assayed in a flask of sintable size, and proceed by one of the methods

given below isi (a) For dry or semi-dry materials that contain no appreciable amount of basic substances Add a volume of I N sulfure acid equal in ml to not less than ten times the dry weight of the material in g , but the resulting solution shall contain not more than 50 mg of meeting acid per inl If the material is not readily soluble, comminute it so that it may be evenly dispersed in the liquid, then agitate vigorously and wash down the sides of the flask with I \ sulfuric acid

Heat the mixture in an autoclave at 121° to 123° for 30 minutes and cool 1f lumping occurs, agitate the mixture until the particles are exculy dispersed. Adjust the mixture to a pll of 68 with sodium hydroxide solution, dilute with water to make a final measured volume that contains approximately 0.1 µg of nicotinic acid in each ml. and filter if the solution is not clear

(b) For dry or semi-dry materials that contain appreciable amounts of basic substances Add sufficient sulfuric acid solution to bring the pill of the mixture to between 50 and 60 Add such an amount of water that the total volume of iqui I shall be equal in ml to not less than ten times the dry weight of the sample in g , but the resulting solution at all contain not more than 50 mg of nicotinic acid in each ml Then add the equivalent of 10 ml of 10 \ sulfuric acid for each 100 ml of liquid and proceed as directed under (a), beginning with the second sentence

(c) For liquid materials Adjust the material to a pH of 50 to 60 with either sul furio acid solution or sodium hydroxide solution, and proceed as directed under (b)

beginning with the second sentence STANDARD MICOTINIC ACID STOCK SOLUTION I DISSOlve 50 mg of USP Micotinic Acid Reference Standard, previously dried and stored in the dark in a deaccator over phosphorua pentoxide, in sufficient alcohol to make 200 ml Store in a refrigerator

Each 10 ml represents 100 ug of USP \scotinic Acid Reference Standard STANDARD MICOTINIC ACID STOCK SOLUTION II To 100 ml of atock solution I, add sufficient water to make 1000 ml Store under toluene in a refrigerator Each 10 ml

represents 10 ug of USP \icotinic Acid Reference Standard STANDARD MCOTINIC ACID SOLUTION Dilute 10 ml of stock solution II with suffi cient water to make 1000 ml I ach I 0 ml represents 0 1 µg of USP Nicotinic Acid Reference Standard Prepare fresh standard solution for each assay

BASAL MEDIUM STOCK SOLUTION	1
Acid hydrolyzed Casem Solution	25 ml
Cystine Tryptophan Solution	25 ml
Dextrose, Anhydrous	10 g
Sodium Acetate Anhydrous	5 g
Adenine-Guanine Uracil Solution	5 ml
Riboflavin-Thiamine Biotin Solution	5 ml
p-Aminobensoic Acid Calcium Pantothenate Pyridoxine	
Solution	5 ml
Salt Solution A	5 ml
Salt Solution B	5 m

Dissolve the anhydrous dextrose and sodium acrtate in the solutions previously mixed and adjust to a pH of 68 with sodium by droxide T S [Test Solution] Finally add water to make 2.0 ml

ACID HYDROLTZED CASELY SOLUTION MIX 100 g of vitamin free easein with 500 mi of constant-boiling hydrochloric act I (approximately 20 per cent HCl) and reflux

¹⁹³ The concentrations of the sulfuric send and sodium hydroxide solutions used are not stated in each instance because these concentrations may be varied depending upon the amount of material taken for assay witume of test within and luffering effect of material

the mixture for 24 hours. Remove the hydrochloric neid from the mixture by distillation under reduced pressure until a thick paste remains. Redissolve the resulting paste in water, adjust the solution to n pH of 35 (±0 1) with sodium hydrocude TS, and add water to make 1000 ml. Add 20 g of netwated charcoal, stir for 1 hour, and filter. Repeat the treatment with activated charcoal if the filtrate does not appear straw colored to coloriess. Store under toluene in a refrigerator. Filter the solution of in preputate forms upon storage.

CYSTINE TRIPTOPHAN SOLUTION Suspend 40 g of L-cystine and 10 g of L-tryptophan (or 20 g of D_LL-tryptophan) in 700 to 800 ml of water, heat to 70° to 80° and add 20 per cent by droelilone acid dropwise, with attring, until the solids are dis solved (Cool, and add water to make 1000 ml Store under toluene in a refrigerator at a

temperature not below 10°

AONN'S GUANNE-URACIL SOLUTION Desolve 0.1 g each of ademne sulfate, guanne hydrochiorde, and uracil, with the aid of heat, in 5.0 ml of 20 per cent hydrochlore acid, cool, and add water to make 100 ml Store under toluene in a refrigerator

RHOPLAYIN THAMINE HYDROCHLORIDE BIOTHY SOLUTION Prepare a solution containing in each ml, 20 μ g of rhoflaxin, 10 μ g of thamine by drochloride, and 0 04 μ g of biotin, by dissolving crystalline riboflavin, crystalline thamine hydrochloride, and crystalline biotin (free acid) in 0 02 N neetic acid. Store, protected from light, under toluene in a refrigerator

p-ANTIOBENZOIC ACID-CALCIUM PANTOTHENATE-PYRIDOXINE HYDROCHLORIDE SOLUTION Prepare in solution in neutral 25 per cent alcohol to contain 10 µg of p-aminobenzoic acid, 20 µg of calcium pantothenate, and 40 µg of pyridoxine hydrochloride in each ml. Store in a refuserator

SALT SOLUTION A Dissolve 25 g of monobasic potassium phosphate and 25 g of dibasic potassium phosphate in water to make 500 ml Add 5 drops of hydrochloric and and store under tolures.

Salt Solution B Dissolve 10 g of magnesium sulfate, 05 g of sodium chloride, 05 g of ferrous sulfate, and 05 g of manganese sulfate in water to make 500 ml Add 5 drops of hy drechloric seel and store under toluene

Stock Culture of Lactobactillus arabinosus 17-5 Dissolve 20 g of watersoluble yeast extract in 100 ml of water add 0.5 g of anhydrous dextrose, 0.5 g of anhydrous sodium acctate, and 1.5 g of agar, and heat the inxture, with stirring, on a steam bath, until the agar dissolves Add approximately 10-ml portions of the hot solution to test tubes plug the tubes with cotton, sternlize in an autoclaye at 121° to 123°, and allow the tubes to cool in an upright position Prepare stab cultures in 3 or more of the tubes using a pure culture of Lacobactilus arabinosus 17-5, 121 incubating for 16 to 24 hours at any selected temperature between 30° and 37° but held constant to within ±0.5° and finally store in a refrigerator Prepare a fresh stab, of the stock culture every week and do not use for incoclum if the culture is more than 1 week, old

CULTURE MEDIUM To each of a series of test tubes containing 5.0 ml of the basal medium stock solution, and 5.0 ml, of water containing 1.0 pg of nectinic acid. Plug the tubes with cotton, sternize in an autoclave in 121° to 123°, and cool

INOCULUM Make a transfer of cells from the stock culture of Laclobacillus arabinosus 17-5 to a sterile tube containing 10 ml of culture medium Incubate this culture for 16 to 24 hours at any selected temperature between 30° and 37° but beld constant to within ±0.5° The cell suspension so obtained is the inoculum

Assay Procedure Prepare standard alcotinic acid tubes as follows To duplicate test tubes, add 00 ml, 05 ml, 10 ml, 15 ml, 20 ml, 25 ml, 30 ml, 35 ml, 40 ml, 45 ml, and 50 ml, respectively, of the standard

³³ Pure cultures of Lactobacillis aral mosus 17 5 may be obtained from the American Type Culture Collection 10³⁰ M Street V W Washington C D C as number 8014

tion and water to make 10 ml. Prepare tubes containing the material to be assayed as follows: To duplicate test tubes add, respectively, 1.0 ml., 2.0 ml., 3.0 ml., and 4.0 ml. of the

test solution of the material to be assayed. To each tube add 5.0 ml. of hasal medium stock solution and water to make 10 ml. After mixing, plug the tubes with cotton or cover with caps, and sterilize in an autoclave at 121° to 123, in Cool, aseptically inoculate each tube with I drop of Inoculum, and incubate for 72 bours at any selected temperature between 30° and 37°, but held constant to within ±0.5°. Contamination of the assay tubes with any foreign organism invalidates the assay.

Titrate the contents of each tube with 0.1 N sodium hydroxide, using bromothy mol blue T.S. as the indicator, or to a pli of 6.8 measured electrometrically.

CALCULATION Prepare a standard curve of the nicotinic acid standard titrations by plotting the average of the titration values expressed in ml of 0.1 N sodium by droxide for each level of the meotimic acid standard solution used, against ag of meetinic acid contained in the respective tubes. From this standard curve, determine hy interpolation the meeting and content of the test solution in each tube. Discard any values of more than 0 40 or less than 0 05 pg of meotime acid in each tube Calculate the meetinic acid content in each ml. of the test solution for each of the tubes. The meeting acid content of the test material is calculated from the average of the values obtained from not less than 6 of these tubes that do not vary by more than ±10 per cent from the average If the titration values of less than 6 of these tubes containing the test solution are within the range of the titration values of the meetinic acid stand. ard tubes containing 0.05 to 0.40 µg of meetings and, the data are insufficient to permit calculation of meeting acid content of the test material. Titration values exceeding 10 ml. for the tubes of the standard meeting acid solution series containing 00 ml, of the solution indicate the presence of an excessive amount of meeting acid in the basal medium stock solution and invalidate the assay.

PYRIDOXINE AND RELATED COMPOUNDS (VITAMIN B4)

Goldberger and Lillie in 1926 reported the development of a characteristic dermatitis, called acrodynia, in rats fed a diet deficient in what they regarded as the pellagra-preventive (P-P) factor In 1934, recognizing the multiple nature of the vitamin B complex, P Gyorgy 154 design nated the missing factor vitamin Be, the rat pellagra-preventive factor, the deficiency being characterized by edema and denuding of the paws and the areas around the nove and mouth, and thickening of the ears Lack of this vitamin was also shown to induce microcytic hypochromic anemia and neurologic lesions in dogs and pigs Vitamin Be attracted so much attention that within one year, 1938, it was isolated independently hy five groups—Lepkovsky, Keresztesy and Stevens, P György, Kuhn and Wendt, and Itiba and Miti In the following year, the chemical structure was elucidated and the compound was synthesized independently by Keresztesy, Stevens, Harris, Stiller, and Folkers in the United States and by Kuhn, Westphal, Wendt, and Westphal in Germany The vitamin was named pyridoxine by the American investigators and adermin by the Germans

Overheating of the assay tubes during sterilization may produce unestimation results. 11 Gybrey \ature, 133, 495 (1934)

It should be noted that the term vitamin Be includes not only nyridovine but pyridoxal and pyridoxamine as well, all three compounds being found in noture

This is the reason for the anomalous results obtained in microbiological185 and chemical assays186 for vitamin B. which led to the discovery of the aldehyde and amine derivatives, pyridoxal and pyridoxamine. These derivatives have hiological activity equal to that of pyridoxine for rats and somewhat less for chicks, but their utilization by microorganisms varies considerably. All three compounds are equally active for Saccharomuces carlsbergensis, while pyridoxal and pyridoxamine are less active than pyridoxine for Saccharomyces cervisiae; on the other hand the aldehyde and amine are relatively more active for L. belreticus or S. fecalis B. and only pyridoxal has activity for L. casei. Evidence suggests the existence of members of the vitamin Bs complex as yet unknown. 187 It is probable that these are known forms bound in such a way that they are not released by conventional methods of bydrolysis.

Physiological and Clinical Aspects of Pyridoxine, Deficiency of vitamin Be in rats causes acrodynia (see Fig. 279), edema, inhibition of growth, and nerve degeneration. The vitamin is essential for the chick. rat. dog, and pig, and for numerous microorganisms. The significance of the vitamin is claimed to be related in part to the metabolism of unsaturated fatty acids. The dermatitis observed in vitamin B. deficiency closely resembles that noted in cases of fatty acid deficiency. Some observers have reported that the addition of certain unsaturated fatty acids to a diet deficient in vitamin Be protects against the appearance of the de-

ficiency syndrome of the vitamin

The role of pyridoxine in amino acid metabolism is illustrated by the increased severity of symptoms upon the addition to a pyridoxine-poor diet of supplemental levels of cystine, methionine, tryptophan, glycine, or serine. Pyridoxine-deficient infants188 or rats189 lose their normal capacity to form miscin and N1-methylnicotinamide from tryptophan. However the dependence of the tryptophan conversion on pyridoxine is questioned by some investigators. 190 Vitamin B. deficiency in rats and swine is characterized by the appearance in the urine of xanthurenic acid, a metabolite of tryntonhan. All three known members of the vitamin B, group-pyridoxine. pyridoxal, and pyridoxamine—are converted by organisms capable of utilizing them into amino acid decarboxylases. 191 Pyridoxal phosphate functions as a coenzyme which, when combined with the apoenzyme (the protein moiety of the enzyme), such as is present in Strentococcus fecalis R grown in a medium deficient in vitamia B6, catalyzes

¹¹⁶ Snell: J. Biol. Chem., 157, 491 (1945).

[&]quot;Sincili J. Just. Liet., 194, 494, 13849).
"Hochberg, Melnick, and Oser: J. Biol. Chem., 184, 313 (1944); 155, 119 (1944),
"Hochberg, Himes, and Oser: J. Biol. Chem., 180, 1 (1945),
"Syndeman, Holt, Carretero, and Jacobs. J. Clin. Nutrition. 1, 200 (1953),
"Syndeman, Holt, Carretero, and Jacobs. J. Clin. Nutrition. 1, 200 (1953).

Schweigert and Pearson: J. Biol. Chem., 168, 555 (1947). 100 Sarett: J. Biol. Chem., 182, 691 (1950).

Heimberg, Rosen, Leder, and Perlzweig: Arch. Biochem., 28, 225 (1950). ¹¹ Gunsalus and Bellamy: J. Biol. Chem. 155, 357 (1914); Rellamy, Umbreit, and Gunsalus: J. Biol. Chem., 160, 461 (1945); O'Kane and Gunsalus: J. Biol. Chem., 170, 425, 433 (1947),

decarboxylation reactions Amino acids reported to require pyridox phosphate as a coenzyme for decarboxylation include tyrosine lysing arginine, ornithine glutamic acid, and dioxyphiciylalanine ("dopa" the end products being respectively, tyramine, cadaverne agriating putrescence, y aminobutyric acid, and 3,4-dibydroxyphenylethylamin



Fig 279 Pyridoxine-deficient rat Note edema and dermatitis of paws and nose

The high specificity of decarboxylation reactions permits their use for inalytical purposes. Glutamic acid decarboxylase is present in the brain issue of rats and in vitamin Br-deficiency it is the pyridoxal phosphate social properties that it is depleted rather than the apoenzyme is.

The same coenzyme pyridoxal phosphate, has been demonstrated to function in transamination reactions¹⁸¹ of which the following is an

²³ For studies of this glutamic send decarbox laws see papers by Awapara & al. Uden friend and Roberts & al in J Bod Chem 19,0-51

²³ Lichtein Gunsalus and Imphret J Bod Chem 161 311 (1945)

example Such reactions are suggested as the explanation of the growth response of rats to keto and hydroxy analogs of certain amino acids 194 Evidence also indicates that vitamin B₆ participates in enzyme systems uncled in the synthesis of amino acids by microorganisms

The fact that pyridovine derivatives play a part in such a variety of amino acid revetions would seem to justify the belief that vitamin B_6 is required in the nutrition of man as it is for rodents, fowl, and other species. While no clear-cut deficiency syndrome is seen chincally which is comparable to thirmine or macin deficiency, vitamin B_6 deficiency has heen produced experimentally in man ¹⁸⁵. The daily human requirement of vitamin B_6 estimated chiefly on the basis of animal experiments, is 2 mg. The vitamin B_6 compounds are excreted mainly as 4-pyridovic acid.

Pyridoune has been used clinically in aene and other dermatologic disorders, but its value has yet to be conclusively demonstrated Parkinson's disease, muscular dystrophy, and especially nausea and vomiting of pregnancy are among the conditions which have been treated with pyridoxide, but the results are too indefinite to justify associating these conditions with vitamin deficiencies, particularly of the B group, generally are of a multiple nature Pellagrins who responded only partially to the administration of thiamine, riboflavin, and macin showed considerable improvement after intravenous administration of pyridoxine.

Storage and Synthesis of Vitamin B₆. Vitamin B₅ is present in most animal tissues, with high concentrations in the liver. It is synthesized by

bacteria in the rumen of sheep and cattle

Distribution of Vitamin B₆ 138 In natural materials vitamin B₆ occurs principally bound to proteins. Rich sources are yeast and rice polishings Seeds and cereals are good sources especially the germ. In rice brain the vitamin is present to a small extent in the free form, but a major portion is a bound complex of pyridovine, readdy hydrolyzable by heating with strong acids. Pyridovine was isolated from rice polishings in 1932 by Ohdake before its identity as a vitamin was known. In yeast and liver, it is present chiefly as bound by ridovamine, though small concentrations of pyridovine and pyridoval are also found.

¹¹⁴ Meister and Wł te J Biol Chem 191 211 (1951) Holden Wildman and Snell Ibi 1 191 509 (1951)

¹⁹³ Mueller and Vilter J Clin Invest 29 193 (1950) 24 See Append x IV

Chemistry of Vitamin B. Pyridoxine hydrochloride is a white, odorless, slightly bitter, crystalline solid, melting at 207° C with slight decomposition It is optically mactive and very soluble in water, slightly soluble in 95 per cent alcohol and in acctone, and insoluble in ether. The free bace is also a colorless, crystalline solid, melting at 160° C. It is soluble in water, acetone, and alcohol, and slightly soluble in other and chloroform The aqueous solution of pyridoxine hydrochloride has a pH of approximately 3 It is adsorbed from acid solution on zeolite, charcoal, or fuller's earth, and may be eluted from zeolite with 10 per cent potassium chloride or from fuller's earth with a weak alkali. The structures of pyridoxine (2methyl-3 hydroxy-4,5bis(hydroxymethyl)pyridine), pyridoxal, and pyridoxamine are

Pyridovine, pyridovamine, and pyridoxal can be separated by paper chromatography and distinguished by their relative positions on the strip For this purpose a "bioautographie" procedure developed in the semor authors' laboratory 157 is used The developed strip is laid flat on an agar plate containing a Be-deficient medium and lightly seeded with Saccharomyces carlsbergensis, an organism capable of utilizing any of these forms of the vitamin After incubation, the loci of Befactors on the strip are recognized by the zones of yeast growth

Early chemical and microbiological studies of vitamin Be were complieated by the presence of the several forms, and by the facility of transformation of one compound to another Conditions favoring amination or partial oxidation of pyridoxine result in the formation of some of the amine or aldehyde This is attended by considerable change in the capac-

ity for growth stimulation of certain microorganisms

The three known forms of vitamin Be possess characteristic absorption curves in the ultraviolet region of the spectrum These curves show no absorption maximum common to all three compounds However, the total concentration of vitamin Be present in a solution relatively free of interfering substances may be determined by measurement at 325 ma and pH 675 Under these conditions, the Eise value for each of the compounds is 110 186

Pyridovine, pyridoxal, and pyridoxamine are destroyed by exposure to light Destruction is most rapid in the ultraviolet region of the spectrum in neutral or alkaline solution Though all three compounds are comparatively stable to light in 0 1 N acid, pyridoxamine is slightly sensitive. The photolysis of vitamin B₀ is not affected by the presence of air All three forms of the vitamin are stable to heating at 100° C with 5 N sulfurn or hydrochloric acid, only pyridoxal is unstable to heat in alkaline solutions All three compounds are stable to mild oxidizing agents like manganese dioxide in alkaline solution or hydrogen peroxide in neutral solution at room temperature. However, they are destroyed by potassium permanganate or by manganese dioxide in acid solution at room temperature, or by heating with nitric acid, potassium permanganate, or hydrogen peroxide.

DETERMINATION OF VITAMIN B6

Since a major proportion of the vitamin B₄ present in natural materials occurs in bound forms, assays must be preceded by hydrolysis by enzymes or by heating with acid. Pyridoxine reacts with ferric chloride to produce a compound having an orange-red color. Pyridoxine is phenolic and reacts with diazonium salts to produce dyes. The other forms probably have similar reactions, Pyridoxine, pyridoxal, and pyridoxamine comple with 2,6-dichloroquinone chloroimide to produce a blue dye, the last two being about half as reactive as the first. The reaction with pyridoxine is shown in the following equation. In the presence of boric acid, pyridoxine alone forms a stable complex which does not couple with the reagent, thus permitting its differentiation from pyridoxal and pyridoxamine in pure systems. ¹²⁵ The low values for the pyridoxine content of natural materials

Pyridoxine

2,6-Dichloroquinone chloroimide

Blue dye

obtained by this procedure indicated the existence of other, possibly conjugated, forms of vitamin B_s. At any rate it is not recommended as an analytical method except in relatively pure mixtures where pyridoxine is the predominant B_s factor, as in pharmaceutical preparations.

Microbiological methods for the determination of vitamin Be are based on the growth stimulation of yeast or baeteria. These are complicated by the variable responses of the microorganisms to the different forms of the vitamin. All three forms have equal activity on a molar basis for Saccharomyces carlsbergensis. For Saccharomyces cerevisiae, however, the amine has 40 per cent, and the aldehyde 46 per cent, of the activity of pyridoxine. Though pyridoxine itself is inactive for Lactobacıllus caser and Streptococcus fecalis R, pyridoxal is active and pyridoxamine inactive for the former, whereas pyridoxamine is active and pyridoxal 36 per cent as active as pyridoxamine for the latter. Assuming that vitamin Be complex contains no other members than pyridoxine, pyridoxal, and pyridoxamine, differential assays may be conducted on natural materials or on mixtures of the three synthetic forms, employing several microorganisms Such assays are difficult, however, because of the ease of conversion from one form to any other. Pyrido al when autoclaved with hydrolyzed casein produces some pyridoxamine, resulting in increased activity for S. fecalis. The reverse of this transamination reaction occurs when pyridoxamine is autoclaved with a-ketoglitarie acid. Such treatment results in increased activity for L cases and decreased activity for S fecalis Autoclaving pyridoxine with eystine, glycine, ammonia, or thioglycollic acid or treating with bydrogen peroxide results in the formation of "pseudopyridoxine" (probably pyridoxal and pyridoxamine), having in creased biological activity for lactic acid organisms. Attempts have been made to prevent these transamination or partial oxidation reactions by sterilizing the hydrolyzed samples and the microbiological media sepa rately, and then combining them a septically Such procedures, bowever, may not give a true picture of the concentrations of the various forms originally present in the test materials, since the undesirable conversions may occur during the sterilization of the samples themselves When animal tissues are autoclaved with pyridoxine, the activity for certain lactic acid bacteria may increase several thousandfold Moreover, the differential assays are based on the assumption that pyridoxine, pyridoxal and pyridoxamine are the only existing forms of vitamin Bs

Certain anomalous results obtained in the assay of natural material for vitamin B. have been interpreted to indicate the presence of forms of the vitamin as yet unknown. The bound pyridoxine in rice brain concentrates is completely liberated by autoclaving with 2 N sulfurin acid, so that analysis of such hydrolyzates with Saccharomyces carlsbergens gives values agreeing with those of the rat assay. Hydrolysis of veast and liver samples by the same treatment, however, gives low values by the merobiological procedure. When the sample is hydrolyzed by heating with comparatively weak acid, 0.055. Sulfurie, the values agree with those found in the rat assay. The low results are not attributable to destruction of the known forms of vitamin B, by the strong acid treat ment, since pyridoxine, pyridoxal, or pyridoxamine may be heated in the presence of yeast and liver without measurable destruction.

Since all members of the vitamin B_c complex, both known and unknown have equal activity for Saccharomyces carlsbergensis and for the rat, reliable microbiological assays of natural materials may be made employing this microorganism. As described above, the concentration of sulfure acid required for the hydrolytic liberation of bound forms of the vitamin varies with the type of product under assay. Hence careful attention must be given to this prehiminary step. To destroy thamner which, under certain conditions may interfere with the pyridovine assay, it has been suggested that the sample be hydrolyzed with \(\) sodium hydrovide under pressure \(\)

Biological procedures have been described for the estimation of vitamin B_t based on the growth of rats and on the cure of specific rat dermattion of these the methods of Dimick and Schreffler's and of Sarma Suell and Flythjem " have been widely used. The latter authors have also reported a chick-growth assay.

The principal urmary exerction product resulting from the ingestion of

hydroxymethyl pyridine). This compound is converted by heating with strong acid to a highly fluorescent lactone which can be determined by simple fluorometric means ²⁰¹ This reaction is useful in metabolic studies of vitamin B₆.

Determination of the Vitamin B. Complex (Microbiological Method of Atkin, Schultz, Williams, and Frey).** Principle. The growth stimulation of a strain of the yeast Saccharomyces carlsbergensis by vitamia B. is employed for the microbiological assay of the vitamin.

Procedurc: Preporotion of Yeost Inoculum. Prepare a fresh slant of Culture 4228 (Amerlean Type Culture Collection), a strain of Socchoromyces carlsbergensis, on "Difco" mait agar and incubate for 24 hours at 30° C. Remove a quantity of fresh growth with a sterile wire loop and suspend in 10 mi. sterile 0.9 per cent saline in a colorimeter tube. With the aid of a densitometer*30 or photoelectric colorimeter, adjust the concentration to an equivalent of 1 mg. moist yeast per ml. by adding sterile saline. Dilute 5 ml. of the adjusted suspension with 45 ml. saline in a sterile Etlenmeyer flask.

Preporotion of the Somple. Suspend a portion of the sample containing between 2 and 4 pg. vitamin Be in 180 ml. 0.055 N sulfuric acid. 304 Autoclave at 15 pounds pressure for 1.5 hours, cool, neutralize to pll 5.2, and dilute to 200 mi. Centrilugo if turbid and assay the clear supernatant extract.

Preporation of Bosol Medium. Mix 100 ml. sugar and salts colution, 102 ml. potassium citrate buffer, 20 ml. casela hydrolyzate, 10 ml. thiamine solution, 10 ml. inositol solution, 4 ml. biotin solution, 5 ml. calcium pantothenare solutioa, and 10 ml. niacin solution. Dilute to 200 ml.

Microbiological Assay. Place 5 mi. basal medium in each of a series of 18-mm. pyrex test tubes. In successive tubes pipet respectively 0.25, 0.50, 1.00, 2.00, 3 00, and 4.00 ml. of extract. With each assay series also include a reference series consisting of tubes containing 0, 5, 10, 15, 20, 30, and 40 millimicrograms pyridoxine hydrochloride. Adjust the total volume in each tube to 9 mi. Plug the tubes and steam for 10 minutes and cool. Under asseptic

¹⁰¹ Huff and Perlawerg J. Biol Chem , 155, 345 (1914).

³⁹ Atkin, Schultz, Williams, and Frey Ind Eng Chem, Anal Ed., 15, 141 (1943). The addition of macin to the basal medium was proposed by Hopkins and Pennington Biochem J. 41, 110 (1947).

²⁰² Calibrate the densitometer with a suspension of moist baker's yeast.

²⁰ N sulfuric acid wheat, and wheat products require more acid For such samples use

Super and Saltz Solution Dissolve 200 g C P destrose (anh) drum), 2.2 g mono-potassum phosphinte, 1.7 g potassum phorode, 0.5 g calcium chloride dhip drate, 0.5 g magnessum sulfate, 0.01 g ferrie chloride, and 0.01 g manganese sulfate in distilled water and dilute to 1000 ml

Polassium Curate Buffer Dissolve 100 g potassium eitrate monohydrate and 20 g citric acid monohydrate in distilled water and dilute to 1 liter

Casein Hydroly ate Solution Neutralize 80 ml "vatamin-free" casein hydrolyzate (10 per cent solution, obtainable from General Biochemicals, Inc., Chagrin Palls, O) to pl14 6 and dilute to 100 ml

Thiamine Solution 10 µg per ml Inositol Solution 1 ing per ml

Biotin Solution 0 S µg per tul

Calcium Pantothenate Solution 200 pg per ml Niacin Solution, 0 1 mg per ml.

conditions introduce into each tube 1 ml. of the yeast inoculum Place the tubes in a mechanical shaker for 16-18 hours at 30°C Immediately thereafter estimate the yeast growth turbulimetrically in a densitometer or photoelectric colormeter with a 660 or 720 ma filter ²⁹⁶

CALCULATION Plot the results of the reference series in per cent absorption against mag, of puriforments of puriforments of puriforments of the sample for each tube. Average all values which agree within 10 per cent of their mean.

Interpretation. The vitamin B₆ group consists of pyridoxine, pyridoxin, pyridoxinine, and one or more unidentified labile factors All members of the complex show comparable activity for Saccharomyct carlistergensis (Culture 4228) and the rat. Hence this procedure measure biological vitamin B₆ activity rather than the concentration of pyridoxine or any single derivative.

indication that it plays a fundamental role in metabolism. The nature of its function has become clear since the discovery by Lipmann and associates²¹² of coenzyme A, of which pantothenic acid is a molecular component. It is now believed that coenzyme A is the functional form of pantothenic acid in plant and animal tissues and in microorganisms.

Pantothenic acid is required by the chick, rat, pig, dog, and other vertebrates. Deficiency symptoms are quite varied in different species. In the chick, the syndrome includes keratitis, dermatitis, fatty liver, lesions of the spinal cord, and involution of the thymus; the vitamin is necessary for reproduction in hens but not for egg production In rats, insufficient intake of dietary pantothenic acid results in necrotic lesions of the adrenal cortex and other related symptoms. Regulation of salt and water balance are functions of the adrenal cortex. Pantothenic acid deficiency results in increased appetite for salt, and a low-salt diet promotes the graying of hair which results from pantothenic acid deficiency. Deprivation of water produces "bloody whiskers" (i.e., porphyrin staining) in rats, a condition also brought about hy pantothenic acid deficiency.

Achromotrichia or graving of hair in black rats ("rusting" in white rats) on diets lacking pantothenic acid, and similar phenomena in dogs and foves, probably are due to a complex deficiency, since pantothenic acid itself is less effective than extracts containing mixtures of B-complex factors. Pantothenic acid has not heen established to he of specific value

in restoring the color to human gray hair.

The "spectacled eyo" condition in rats, associated with a deficiency of inositol or hiotin, may develop in the absence of sufficient pantothenic acid. Pantothenic acid is required in the metabolism of all hacteria. Those which do not require an external supply synthesize it themselves. Dietary deficiency of this factor may result in secondary vitamin deficiencies due to diminished synthesis by intestinal flora.

. The physiologically active form of pantothenic acid in the animal organism appears to be coenzyme A. This coenzyme (abbreviated CoA) participates in a wide variety of reactions involving the transfer of acetate

or two-earbon muts. Its acetyl derivative (acetyl CoA)

CoA + acetate CoA·COCH₃ Acetyl CoA

is considered to be synonymous with "active acetate" and has been described as the "pivot on which acetyl transfer mechanisms tirn."

However, it is necessary to point out that many of these metabolic transformations are quite complex and involve the mediation of numerous enzymes and coenzymes. CoA participates in the acetylation of amines; examples are sulfanilamide, glycine, etc. In these reactions the carbonyl group of the acetyl radical is involved:

$CoA \cdot C*OCH_3 + R \cdot NH_2 \rightarrow R \cdot NH \cdot C*OCH_3 + CoA$

Lipmann, Kaplan, Novelli, Tuttle, and Guirard J. Biol. Chem., 167, 869 (1947);
 186, 215 (1989).
 Welch and Nichol: Ann. Rev. Biochem., 21, 633 (1952).

In contrast with this mechanism, the methyl group is involved in CoA condensation reactions such as the formation of citric from oxalacetic acid

a reaction mediated by a condensing enzyme *14

It is believed that all substances (e.g., acetaldeliy de, acetate) entenng the tricarbovylic acid cycle as two-carbon fragments generate acetyl CoA 215 This may be illustrated by the following reactions for acetate and pyruvate

The foregoing indicates, in part at least, the vital role played by the pantothenic acid coenzyme in carbohydrate metabolism Acetyl transfer also occurs in the metabolism of fatty acids and β keto fatty acids In ritro studies on butyrate oxidation216 have suggested that for every two earbons of a fatty acid chain one mole of acetyl CoA is generated, in a secondary reaction acetyl phosphate is formed releasing CoA for further degradation of the chain It has furthermore been demonstrated that two moles of acetyl phosphate can be condensed to form acetoacetate What relation these facts have to the biosynthesis of fatty acid is not yet established since labeled acetoacetic acid is not incorporated into fatty acid in rat hver preparations although labeled cholesterol is formed

The role of two-carbon fragments in intermediary metabolism is under active investigation and reference should be made to Chapter 33 and to the current literature (see Bihliography, p 1293) Much of the work in this field is based on in vitro studies of microorganisms, tissue slices, and ex tracts thereof, and its significance for the living animal awaits further study

The bound forms of pantotheme acid in natural materials are available to higher animals but not to microorgamisms. For the latter, it must first be released by suitable hydrolytic procedures Certain synthetic forms of bound pantothenate have been investigated Ethyl monoacetylpantothenate is active for the rat and chick, and ethyl pantothenate for the rat However, neither of these forms shows activity for microorganisms unless they are previously hydrolyzed by procedures similar to the employed for the release of bound pantothenate in natural materials

Certain structurally related compounds behave as antivitamins to ward pantothenic acid Pantoy Itaurine (the sulfonic analog), pantoy

²¹⁴ Ochoa Stern and Schneider J Biol Chem 193 691 (1951)
² Ochoa and Stern Ann Per Biochem 21 547 (1952)

^{1 *} Barker in I hosphorus Metabolism Baltimore Johns Hopkins University Press 1951

tauramine, and homopantoyltaurine inhibit the growth of bacteria which require an external source of the vitamin. These autivitamins probably operate by competing in some enzyme systems. One observer has found that the administration of pantoyltaurine to mice produces symptoms similar to those of pantothenic and deficiency. This observation could not be confirmed by other investigators. For a further discussion of anti-vitamins, see Chanter 36.

Possibly because of intestinal synthesis, pantothenic acid deficiency has not heen observed in man and hence its importance in human nutrition has not been clanfied. The duily requirement, as estimated from the analysis of good diets and from urinary everetion data is approximately 5-10 mg, per 2500 calories. Pantothenic and is present in normal human.

blood or plasma to the extent of 20 to 40 µg per 100 ml

The physiological effect of pantotheme acid is highly specific for that molecule. Replacement of the β alaniue portion with α alanine, or with other amino acids having structures similar to β alanine, produces inactive compounds. The optical antipode of the natural 1-isomer is also matrive. The β -alanine portion alone is sufficient for some yeasts and diphtheria bacilli, and is partly available for rats, but not for chicks. These organisms probably utilize the β alanine for the synthesis of pantothemic acid. Certain hemolytic bacteria require only the dihydroxydimethylbutyryl portion of the molecule. Hydroxy pantothemic acid has a biological activity varying from 2 to 25 per cent of that of pantothemic acid, depending upon the organism employed and the conditions of assay

Storage and Synthesis of Pantothenic Acid. Pantothenic acid from external sources is not required by sheep and cattle, since the vitamin is synthesized by microorganisms in the rumen of these animals. It may be synthesized by certain molds, bacteria, and yeast, when grown in media devoid of pantothenic acid. Some veasts require an external supply of β alumine. Pantothenic acid is produced by green plants after they have developed sufficiently to perform their photosynthetic functions

Distribution of Pantothenic Acid. Pantothenic acid is present in all lung tissues Excellent sources are liver, kidney, rice bran, molasses, egg yolk, peanuts, and peas Appreciable concentrations are also found in sweet potatocs, cats wheat, rye, barley, and broecob Pantothenic acid occurs in natural materials both free and combined, though a major fraction of the vitamin in animal sources is bound While it has not been definitely established that all forms of conjugated pantothenic acid released by un cray matic digestion procedure are equally available for animal species, rat-growth studies indicate this to be so for conzyme A and L bulgarieus factor. In certain foods or extracts (liver powder, yeast, wheat-bran and rice-bran extracts) where rat assays were reported to be considerably higher than microbiological assays, the discrepancies may have been due to unidentified dietary factors or incomplete enzymatic release of pantothenic acid prior to microbiological assays.

Chemistry of Pantothenic Acid. Puntothenic acid (p-)-(a,y-dih) droy,-\(\theta_0\), dimethyl but \(\text{r_j}\))-\(\theta_0\) alanine is a condensation product of \(\theta_0\) alanine and a hydroxyl and methyl substituted but yre acid it is

pale yellow, unstable viscous od

The natural or vitamin-active form of pantothenic acid is the devirorotatory isomer, specific rotation [a]²² = +37.5° Synthetic pantothenic acid is produced commercially as the calcium or sodium salts, in either dextrorotatory or racemic form, the latter having half the potency of the former The dextrorotatory alcohol corresponding to pantothenic acid, pantothenyl alcohol, is vitamin active and is also available commercially

PANTOTHENIC ACID CONTENT OF FOODS (AFTER NEILANDS AND STRONG*)

(AFIER HEILER)	
Food	Pantothense Acid mg per 100 g
Alfalfa	6 2
Beef, dried	5 6 5 5
Egg fresh Laver powder	39 3
Mackerel	1 4 2 6
Salmon Sardines	1 5
Spinach Tuna	2 7 1 4
Wheat germ meal	29

*Neilands and Strong Arch Buckers 19 287 (1948) Based on mucrobiological assistant digestion with liver enzyme and phosphatase

It is claimed to be more stable than the acid at pH below 6 Pantothenic acid is adsorbed by "Nont" at pH 3 5 or hy aluminum ocide previously activated with hydrochlone acid but not by fuller's earth It may be cluted from "Nont" with ammonia or pyridine and methanol Pantothenic acid is soluble in water, ethyl acetate, dioxane, and glacial acetic

acid, and is slightly soluble in ether and amyl alcohol, and insoluble in chloroform and benzene. It is unstable to heat in the dry state or in acid or alkaline medium. In the pH range 5 to 7 it may be autoclaved at 15 pounds pressure for one-half hour without loss.

The hutyrolactone morety of pantotheme acid is known as pantolactone and the corresponding acid is pantone acid, HOCH₂C(CH₃)₂CHOH-COOH A naturally occurring growth factor for certain microorganisms (Lactobacillus bulgaris factor or LBF) has been identified as a \$\textit{\textit{BF}}\$ mercaptothy lamine derivative of removibles and \$\textit{BF}\$ has been identified as a \$\textit{\textit{BF}}\$ mercapto-

eth lamine derivative of pantothenic acid and given the name pantethene. The corresponding disulfide, pantethine, exist presumably in equilibrium with the reduced form. The following diagram illustrates the relationship between these compounds in terms of the various moieties in the molecule of acetyl coenzyme A.

DETERMINATION OF PANTOTHENIC ACID

No satisfactory chemical method is available for the determination of pantotheme acid in natural products, although several procedures have been described for pharmaceutical preparations $One^{2i\tau}$ is based on the hydrolytic release of the β alanine moiety, followed by colorimetric estimation as the alkaline 2,4-dimitrophenylhydrazone. Interfering B vitamins may be removed chromatographically, but ascorbic acid still presents a problem

Another procedure 18 is based on a reaction between the pantoyl lactone mosety and hydroxylamine in sikaline solution and the develop ment of a purple color upon acidification and addition of ferne chloride. In contrast to the previous method this may be applied to the estimation of pantothonyl alcohol, known commercially as "Panthenol' However in terference due to ascorbic acid, pantoic acid, and the lactone must be climinated by adsorption on a basic resum XE-75 18

Determination of Pantothenic Acid (Microbiological). The absolute requirement of a number of microorganisms for free pantothenic acid was primarily responsible for the original discovery of this vitamin Many determinations of the pantothenic acid content of natural materials bave heen published. By far the majority of these have been micro hiological procedures although a considerable amount of work has also heen reported based on the use of chicks,*** rats, and other species as test animals. The repeated observation that animal assays in general gave higher values than the microbial tests, together with several early reports on the release of additional increments of pantothemic acid activity by autoly sis of fissues or treatment with specific enzy me preparations, led to the recognition of "bound forms" of the vitamin which occurred in many natural materials However, the release of activity from these bypothetical conjugates was neither striking nor consistent.

The discovery that coenzy me A was in reality a pantothenate-containing prosthetic group of enzymes responsible for transacety lation in living cells proved to lie the key to the problem of bound or conjugated forms of the vitamin which had been shown to be unavailable to microorganisms such as L arabinosus A specially prepared enzyme obtuined from pigeon or chicken liver was found capable of mactivating coenzymic A with the

²¹⁷ Szalkowski Mader and Frediani Anal Chem 22 369 (1950)
2 1 Wollish and Schmall Anal Chem 22 1033 (1950)

m Gyorgy Rub n and de R tter in I stamm Methods Vol 2 New York Academic Press inc 1951 This res n is available at Res nous Products D v n Rol m and Haas Co Piladelpl a 5 Pa

¹⁷⁴ Jukes J Biol Chem. 117 11 (1937) J Astrition 21 193 (1941)

concurrent liberation of free pantothenate At the same time, an alkaline phosphatase from intestine was shown to split CoA in another manner with the liberation of phosphate 221

By combining in a single treatment both an intestinal phosphatase and the CoA-splitting enzyme from pigeon liver, Neilands and Strong" showed that some natural materials contained as much as four times the amount of pantothenate which could be demonstrated by simple direct assay They also showed that other nonspecific phosphatases were rela tively ineffective in liberating the vitamin On the basis of this work, it is evident that a large part of the data previously accumulated on the dis tribution of pantothenate in foods and tissues is in error and needs rein vestigation Some of the new values reported are given in the table, p 1190

Determination of Pantothenic Acid Modified Microbiological Method 0 Skeggs and Wright 11 Pantothenic acid is determined by measurement of the growth simulation of Lactobacillus arabinosus 17 5 by titration of the lactic and formed or by turi idimetric determination of the cell population. The bound vitamin is first liberated by digestion with alkaline phosphatase and chicken liver enzyme

Procedure Preparation of Iner enzyme 111 Liver from freshly killed chicken or pigeon is chilied, minced, and homogenized with 20 volumes of ice cold acetone in a Waring blendor The acetone insoluble material is filtered and washed with acetone and ether and dried quickly in a current of dry air The resulting liver powder's may be preserved in small, well sealed vials under refrigeration for a considerable period For use, the acetone powder is rubbed in a chilled mortar to a smooth paste with 10 times its weight of cold 0 02 h sodium bicarbonate solution The suspension is centrifuged below 5° C for 30 minutes at 3,000 r p m, and the reddish brown supernatant liquid is pre served by freezing and holding below -20° C

This solution possesses good activity on coenzyme A but has the disadvantage of containing considerable amounts of pantothenic acid in a free form its use in the assay of materials of low pantothenate content presents the prob lem of an excessive blank The amount of free vitamin may be substantially reduced by treatment of the solution with an anion exchange resin no The following procedure is recommended

Prepare the resin for use by washing twice with 10 volumes of N liCl and centrifuging. The acid treated resin is then washed 8 to 10 times with 10 volumes of water until the pll of the wash is about 5 The resin is left in a slurry of a consistency which can be pipetted with a 10 mi serological pipet To I volume of resin slurry is added a few drops of M tris (hydroxymetbyl) aminomethane buffer, pil 8 3, to bring the pH to 8 8, and 1 volume of ice cold liver enzyme solution is added and the suspension stirred for 3 to 5 minutes in an ice bath The suspension is centrifuged at 5 000 r p m. The enzyme solu

III Lapmann Kaplan Novelh Tuttle and Gurard J Biol Chem 167, 869 (1947) Novelli et al J Biol Chem 177 97 (1949) 192 181 (1951)

^{111 \}clambde lands and Strong Arch Blockem 19 287 (1948) sumilar method has been adopted 111 Sheggs and Wight J Blot Chem 15c 21 (1944) A sumilar method has been adopted 115 the Association of Official Agricultural Chemistry J Assoc Official Agr Chemistr 35 by the Association of Official Agricultural Chemistry J Assoc Official Agr 103 (1952)

¹¹¹ Novella and Schmetz J Bul Chem 192 181 (1951)

The prepared acctone powder may be obtained from Armour and Co Chicago III 124 Dowex 1 200-400 mesh obtainable from the Dow Chemical Co Midland Michigan

tion is again treated with another volume of nnion resin slurry and centrifuged. This process usually reduces the free pantothenate content of the solution to less than one-tenth its original value. Preserve the preparation at -20° C, until ready for use.

Preparation of Intestinal Phosphatase. This preparation can be obtained by following the procedure of Schmidt and Thannhauser*** but is more conveniently obtainable from commercial sources.*** A 2 per cent solution of this material usually contains about 100 Schmidt-Thannhauser units per ml, and is quite low in free pantothenic acid activity.

Liberation of Bound Pantothenate. To a sample containing 5 to 15 µg, bound pantothenate are added 0.1 ml. intestinal phosphatase, 0.2 ml. liver enzyme, 0.1 ml. M tris(hydroxymethyl)aminomethane buffer, pH 8.3, and the whole is made to a total volume of 1 ml. Suitable blank and control tubes should also be set up to allow a correction for any free pantothenate activity in the enzyme preparations. The tubes are incubated at 37° C. for 3 hours and then immersed in a boiling water bath to stop enzyme reactions. The samples are then diluted to bring them into the range of the usual standard curve employed in the pantothenic acid assay of Skeggs and Wright described below.

Preparation of Medium. Prepare the basal medium baving the composition shown in the table below. Five liters may be prepared as a stock solution at one time provided the glucose and synthetic vitamins are omitted. This solution keeps indefinitely at room temperature under benzene even witbout sterilization. Prepare a stock vitamin solution containing 4 mg, each of thiamine, riboflavin, and niacin, 8 mg, pyridoxine bydrocbloride, 0.4 mg, p-aminobenzole acid and 10 µg, biotin per 100 ml. Store the vitamin supplement in a dark bottle in the refrigerator and renew monthly. Prepare a solution containing 100 µg, per ml. of calcium pantotbenate.

Culture. Carry stab cultures¹¹⁰ of Lactobacillus orobinosus 17-5 by montbly transfer in a medium containing 1 per cent yeast extract, 1 per cent glucose, and i.5 per cent agar. After transfer, incubate the culture at 33° C, for 24 to 48 hours, then store in the refrigerator. Prepare the inoculum for the assay tubes by transferring from the stock culture to a sterile tube containing 10 ml. of the basal medium to which $0.2~\mu g$. calcium pantothenate has been idded, incubate for 24 hours at 33° C, centrifuge, and discard the supernatant ilquid. Resuspend the cells in 10 ml. physiological saline, centrifuge, and discard the supernatant. Resuspend the cells in sufficient physiological late to produce a very light suspension.

Assay. Into a series of test tubes, piper the following volumes: 0.00, 0.25, 0.50, 0.75, 1.00, 1.25, 1.50, 1.75, and 5.00 ml. Similarly prepare a standard series of tubes from a dilution of the standard solution of calcium pantothenate containing $0.1~\mu\text{g}$, per ml. Dilute the aliquots in both the standard and unknown series with distilled water so that each tube contains a total volume of 5 ml. To each tube add 5 ml. of the double strength medium, cover with cotton plugs or metal caps and heat In an autoclave at 15 pounds pres-

¹¹⁷ Schmidt and Thannhauser J. Biol Chem., 149, 369 (1943) in Intestinal Phosphatase (Calf), Armour and Company, Chicago, ill

²² A culture of the organism may be obtained from the American Type Culture Collection, 2029 M Street, N W, Washington 6, D C, under the classification number 8014

oure for 15 minutes. Add to each tube aseptically one drop of the inoculum prepared as described above. Incubate at 33° C. for 72 hours and determine the lactic acid produced by titration with 01 N sodium hydroxide, employing bromothymol blue as the indicator.

BASAL MEGILM

Ingredient	l er 100 ml Vedium (Double Strength)	Ingre hent	Per 100 ml Medium (Double Strength)	
Cascin ¹²⁸ C, stine Tryptophan Oleic acid Sodium acetate (anh) drous) Adenine Guanne Uracil \anthine Inorganic salts A (see p. 1160)	1 0 g 29 0 mg 20 0 10 0 1 2 g 1 0 mg 1 0 1 0	Inorganic salts B (see p. 1169) Cliucose Thuamine chloride Rhodravin Neotinic acid Pyridoxine hy drochloride p-Aminobenzoic acid Botin pH adjusted to 6 6-6 8	1 0 ml. 4 0 g 200 µg 200 200 400 20 0 0 5	

Equally satisfactory results may be obtained by measurement of the population of the microorganisms turbidimetrically after 18 bours of incuments at 33°C. A photoelectric colorimeter is employed for this purpose The instrument is set at 100 per cent transmittancy with the solution from the assay tube containing no added pantothenic acid.

CALCULATION Prepare a standard reference curve, employing the data obtained in the standard series by plotting on ordinary graph paper, µg of calcium paniothenate as the abscisse and mil of 0 1 \ sodium hydroxide, or photometric density, as the ordinates From this reference curve estimate the calcium paniothenate confent of each tube in the unknown series Divide this value by the volume of the extract originally placed in each tube to order to obtain the concentration per mil of extract Determine the average of at least four values which differ from each other by no more than 10 per cent. Calculate the paniothenate and content of sample using the formula

$$\mathcal{C} \times \frac{D}{G} \times 0$$
92 = $\mu \mathrm{g}$ pantothenic acid per g_of sample

where C is the concentration of calcium pantothenate per ml. of extract D is the final volume to which the extract was diluted G is the weight of the sample taken for analysis and 0.92 is the factor for converting calcium pantothenate to pantothena

Comment. The microbiological determination of pantothenic acid in these enzymic digests employing Lactobacillus arabinosus 17.5 gries values which are in agreement with microbiological procedures employing other microorganisms as well as with the biological chick assay

¹²⁸ Hi drochloric acid hydrolyzed Norit treated vitamin free casem is employed. A 10 per cent solution of casem hydrolyzate suitable for this medium may be obtained from General B or elements In Chagrin Falls Ohio.

PTEROYLGLUTAMIC ACID (FOLIC ACID)

In early investigations, this vitamin was described under various names including vitamia M. factor U. veast Nort eluate factor, vitamin B. and L. cases factor The multiplicity of the nomenclature was oceasigned by the fact that various conjugates of pteroylglutamic acid exist in natural materials. These conjugates vary in their physical and chemical properties, and in their higlogical potencies

Physiological and Clinical Aspects of Pterovlelutamic Acid. This vitamin is a growth factor for various hacteria. Its deficiency in chicks leads to slow growth, poor feathering, and macrocytic anemia In rats a deficiency of pteroylglutamic acid may he produced by addiag sulfonamides to purified dicts Agemia, leukopenia and agranulocytosis are produced and may be cured by administering pteroylelutamic acid Monkeys on a purified diet develop a syndrome characterized by leukopenia, anemia, necrosis of the gums, loss of appetite, diarrhea, and eventual death. The syndrome responds either to pteroviglutamic acid or to pterox liriglutamic acid or to citrovorum factor

Pteroviglutamic acid produces a clinical response in the megaloblastic anemias It is used in the treatment of sprue, nutritional megaloblastic anemia, and the megaloblastic anemias of pregnancy and infancy. The blood picture in these diseases is marked by a low erythrocyte count, reduced hemoglobin, high color and volume indices of the erythrocytes. and lowered leukocytic and platelet counts Accompanying this is "mcgalohlastic arrest" in the hone marrow so that an increase of nucleated red cells, including more than 2 per cent megalohlasts, is seen in films prepared from fluid aspirated from sternal hone marrow Pernicious anemia is included in the megalohlastic group, but the disease is accompanied by neurological changes which respond only to vitamia Bi. In sprue, the administration of pteroylglutamic acid either parentally or orally results first in the disappearance of the symptoms of glossitis A reticulocyte peak is commonly reached in six to mine days. Increases occur in erythrocyte count, per cent hemoglohin, white cell count, and number of platelets in a manner similar to that described above for pernicious agemia. There is an improvement in the sense of well being an increase in appetite, a subsidence of diarrhea, and a gaia in body weight. The hone marrow picture shows a disappearance of the more primative red blood cells and a return of the differential white cell count to normal proportions

Nutritional anemia and the macrocytic anemias of pregnancy and infancy show hemopoletic responses similar to those described for pernicious

anemia and sprue

Pteroylghitamic acid appears to be specific in the treatment of the megaloblastic anemia of pregnancy Typical findings were described in a case by Day and co-workers," and are illustrated in Fig 280 The patient, a parturient woman, had a hemoglobin of 6 5 g per cent and a red cell count of 17 million. The administration of liver extract and vitamin B1. did not relieve the anemia and appeared to aggravate the associated symptoms which included glossitis and diarrhea. The changes following

¹¹¹ Day Hall and Pease Proc Staff Meetings Mayo Clime 24 119 (1919)

administration of pteroylglutamic acid were accompanied by hunger, recession of edema, and disappearance of the other signs and symptoms, including return of the hemopoietic system to normal The increased dietary need for pteroylglutamic and in pregnancy is also manifested in "lactation leukopenia"222 in rats The rats show blood changes due to a deficiency of pteroylglutamic acid in diets which suffice for maintenance but do not meet the extra demand occasioned by inctation

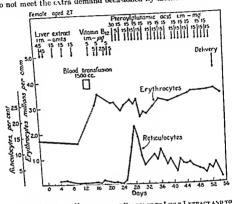


Fig 280 Abspace of Hematopoletic Response to I ivep Extract and to VITAMIN BIE FOLLOWED BY PROMPT RESIGNATION PTEROYICI UTAMIC ACID Courtery Drs Day Hall and I case

Relation of Pteroyigiutamic Acid to Vitamin B₁₂ These two anti anemic substances show some unique clinical interrelationships The blood and bone marrow changes in permicious anemia respond to either pteroylglutamic acid or vitamin Biz even though no dietary deficiencies of pteroylglutamic acid may exist. The uptake of pteroylglutamic acid from the digestive tract of patients with pernicious anemia appears to be normal, but these patients cannot utilize vitamin B12 adequately owing to a functional deficiency of ' intrinsic factor " The megaloblastic anemia of infancy responds uniformly to pteroylglutamie acid but does not respond consistently to vitamin B12 These clinical phenomena find some interesting analogies in biochemistry, for example, both pteroylglutamic acid and vitamin B12 are concerned in biological reactions involving the formation of methionine from homocystine 222

Distribution of Pteroylglutamic Acid Pteroylglutamic acid and its close relative, the citrovorum factor are widely distributed in biological

¹¹¹ Selson and Evans J Autration 38 11 (1919)

¹¹¹ Bennett Science 110 589 (1919)

materials, often as polyglutamic acid conjugates. Good dietary sources include green leaves, asparagus, liver, yeast, cowpeas, and soybeans. The distribution of "folie acid activity" in foods has been exhaustively studied by microbiological and chick assays "14. The name "folic acid" was originally defined in terms of a factor in spinach showing biological activity for Streptococcus fecalis. This factor was probably a mixture of pteroylglutamic acid and "citro original activity for S fecalis and are both present in spinach and many other natural materials.

THE FOLIC ACID ACTIVITY OF FOODS (UNDRIED)

Food	Ug per Kg	Food	Mg per Kg
\pples	0 01	Greens various types	0211
Asparagus	09-14	Lamb, leg	0 03
Barley	0.40	Lamb liver	28
Beans, dried	1-2	Lettuce head	0 03-0 11
' green	01-04	Milk	0 06
' hma	01-06	Oats	0 24
Beef Lidney	0 6	Oranges	0 05
" liver	28-31	Peas	01-04
" round	0 06-0 17	Pork liver	2 2
Brewer's dried 3 cast	20	" loin	0 03
Broccolt	02-15	Potatoes	0 02-0 10
Cabbage	01-07	Raisids	0 01
Cheese, cheddar	0 2 -0 3	Rice, brown	01-03
" cottage	02-03	Soy bears	1 4 -2 7
Chicken liver	3 8	Squasb	0 02-0 24
" meat	0 03	Strawberries	0 05
Corn	0 23	Sweet potatoes	0 05-0 19
Cowpeas	3 3 -4 8	Tomatoes	0 01-0 16
Egg white	0 004	Turkey meat	0 1
" yolk	0 13	Turnips	0 04
		teed II	03-04

* Condensed from Agricultural Handbook No 29 US Dept of Agriculture

Chemistry of Pteroylglutamic Acid. The structure and synthesis of pteroylglutamic acid were described in 1946 *** The molecule consists of a pterioyl group connected in methylene linkage through p-amnobenzou acid to L(+)-vilutamic acid and the structural formula is as follows:

Folic Acid Content of Foods (Agricultural Handbook Vo 29) Washington D. C
 U S Government Printing Office September 1951
 Inger et al., Science 102, 227 (1945), 103 007 (1946)

ne pteridyl group is chemically related to vanthopterin, a yellow pigent which is present in certain natural materials including urine and

e wings of butterflies

Pteroylglutamic acid is a substance of low toxicity as measured by xicological tests with mice, rats, guinea pigs, rabbits, cats, and dogs The Die for mice and rats is about 0.5 g per kg body weight when injected travenously

A second form of vitamin, "fermentation L case: factor," isolated from n aerobic fermentation of an organism of the genus Corynebacterium,



FIG 281 SYNTHETIC PTEROLIGICTANIC ACID CRISTALS, VIEWED THEOLOGIC CROSSED POLABIZERS

Photographed by Dr A. F Kickpatrick Stamford Laboratones, American Cyanamid Co Courtesy Dr T H, Jukes.

has been identified as pteroyldi-y-glutamylglutamic acid. Upon partial degradation with alkali, it yields ni-pteroylglutamic acid "Fermentation L cases factor" is relatively mactive as a growth factor for S fecalis R by comparison with pteroylglutamic acid. However, on a molar basis, it appears to be just as active as pteroylglutamic acid for L cases, for chicks, for rats, and for monkeys When the p-aminobenzoyl-glutamic-acid radical in the pteroylglutamic acid molecule is replaced by p-aminobenzoic acid, the resultant substance is termed "pteroic acid" This substance is active for & fecalis R but not for L cases or for animals

Other members of the pteroylglutamic acid family of vitamins are pteroylhexa-y-glutamy iglutamic acid (previously known as the vitamin B. conjugate), N-formyl pteroic acid (rhizopterin or S lactis R factor) which is mactive for animals, and foliuse acid (citrovorum factor, or leucovorin), discussed in the next section. The differences in biological potency among the various natural forms of pteroylglutamic acid raises a question concerning the significance of microbiological assays in which the preliminary digestion procedures liberate maximum amounts of the parent compound.

Pteroylglutamic acid is a yellow, tasteless substance which is only slightly soluble in water, about 0.01 mg per ml dissolves at room temperature. It is destroyed rapidly by boiling with dilute hydrochloric acid. Its sodium salt is fairly soluble. Light bas a destructive action on the solution. Pteroylglutamic acid bas an absorption spectrum which shows characteristic maxima in the ultraviolet range. The substance crystallizes from water in the form of this lenticular crystals (Fig. 281) which exhibit birefringence and parallel extinction.

DETERMINATION OF PTEROYLGLUTAMIC ACID

Microbiological Assay 237 The assay is carried out with either Laciobacillus cases (ATCC #7469) or Streptococcus fecalis (ATCC #8043)

PTEROYLGUTAMIC ACID STANDARD SOLUTION

(a) Preparation Wash 20 mg of pteroylglutamic acid with water into a 100-ml volumetric flask Add 2 ml 01 N AgOH Shake until material is in solution Add 5 ml of 10 M sodium phosphate buffer, pH 70, and 25 ml absolute ethanol and make up to 100 ml with water

Basal Medium for Assay of Pteroylglutamic Acid with Laclobacillus cases 228

Constituent	Amount per 500 ml double strength medium	Constituent	Amount per 500 ml double- strength medium
	g		mg
Acid by drolyzed casein	5 0	Adenine sulfate	5 0
Sodium acetate	20 0	Guanine hydrochloride	50
Glucose	20 0	Uracıl	5 0
Asparagine	0 30	\anthine	10 0
Tryptophan	0 10	Glutathione	2 5
Cysteine	0 25	Thiamine hydrochloride,	0 2
Tween 80 (219)	0 05	Riboflavin	0.5
K ₂ HPO ₄	0 50	Calcium pantothenate	0 4
KH ₂ PO ₄	0 50	Nicotinic acid	0 4
MgSO,7H O	0 20	Pyridoxine hydrochlor	
NaCl	0 01	ıde	2 0
FeSO, 7H O	0 01	p-Aminobenzoic acid	0.5
MnSO, HtO	0 10	Biotin	0 01

Glucore cysteine and glutathione are added as solids. Other ingredients are added in the form of solutions. Combine ingredients adjust to pH 68 and make up to 500 ml. with water

²³⁴ Stokstad J Biol Chem 149 573 (1943)

²³⁷ Flynn Will ams O Dell and Hogan Anal Chem 23 180 (1951)

²² For assay with Streplococci's feedles om't sodium acetate and KII, PO. Add 27 5 g sodium citrate (dih) drate) and change amount of KrHPO. to 3 10 g

²³ Obtained from Atlas Powder Company Wilmington Del

- (b) Dilutions Dilute to 0 001 μg per ml for assay with L cases; to 0 005 μg per ml for assay with S fecalis
- (c) Assay Levels of Diluted Standard Solutions 01, 02, 04, 06, 08, 10, 15, 20, 30, 50 ml for L cases, 01, 02, 04, 06, 08, 10, 15, 20, 30 ml for S fecalis

The preparation of the various constituents of the medium is as follows:

VITAMIN-FREE, ACID-HYDROLIZED CASEN ** Laboo "witamin-free" casein (300 g) is refluxed at least 12 hours with 3 liters 20 per cent HCl 11 is evaporated under acumum to a thick syrup, then diduted to 3 liters with distilled water. This concentration procedure is repeated twice. The final concentrate is diluted to 1 liter. The pII is adjusted to 2.5 with AOH. The mixture is stirred with 30 g activated carbon (Darco 600) for 15 minutes, then filtered with the aid of "Filter-cel". The pII is adjusted to 6.8 with KOH after the volume of solution is brought to 3 liters, and the charcoal treatment and filtration are repeated. One ml of the solution contains the equivalent of 100 mg original casein.

ADENINE AND GUANINE These are dissolved together in a small amount of 10 N HCl with heat and diluted to 1 mg per ml

XANTHINE AND URACIL. These are dissolved in ddute NH OH and dduted to 1 mg per ml

TRIPTOPHAN Dissolve in a small amount of 10 N HCl and dilute to 10 mg per ml ASPARAGINE One g asparagine is dissolved in 100 ml water

Vitalit's Four mg thismine hydrochlonde, 10 mg riboflavin, 8 mg calcium pantothenate, 8 mg nicotinia acid, 40 mg pyrhdonine hydrochloride, 10 mg paminobenzioe acid, and 200 ng biotin size dissolved in 100 ml distilled water and stored in the cold room under toluene. One ml. of this solution is added per 100 ml. double-threath medium.

bidimeter or after 72 hours' incubation by titration of the acid formed with 0.1 N NaOii. The growth response with *L. casci* may be determined turbidimetrically after 40 hours' incubation, or after 72 hours' incubation by titration of the acid formed with 0.1 N NaOii. From these data a curve can be plotted relating growth to pteroyigiutamic acid concentration, and from this response curve the potency of the unknown samples can be calculated.

CITROVORUM FACTOR (CF)

The occurrence of a growth factor for Leuconostoc citrororum 8081 in certain natural materials including liver extract and yeast was reported in 1948. 242 The organism was shown to respond either to thymidine or to an unidentified "citrororum factor" which was related to ptcroylgiutamic

acid as indicated by the fact that a delayed and submaximal response was produced in the organism by massive doses of pterovlglutamic acid. It was shown that natural materials containing CF would reverse the inhihitory effect of 4aminonterovialutamic acid (aminonterin) for Leuconostoe citrovorum and that the administration of large doses of pteroylglutamic acid to animals produced many-fold increases in urinary CF. The factor was eventually synthesized by the formulation and reduction of oterovigintamic acid

Chemistry of Citrovorum Factor (Leucovorin, Folinic



Fig 282 Calcium Leucovorin, Courtery, Dr T H. Jules American Cianamid Co

Acid). It was shown that 5-formyl-5, 6, 7, 8-tetrahydropteroylglutamic acid had the biological properties of the naturally occurring CF. The compound was synthesized from pteroylglutamic acid by hydrogenation in formic acid, alkaline treatment, and isolation of the active material ²⁴² A photomicrograph of crystals of the calcium salt of leucovorin is shown in Fig. 282

Leucovorin produces responses in the megaloblastic anemias similar to those obtained with pteroylglutamic acid. It blocks the toxic effects of 4-aminopteroylglutamic acid and other "folic acid antagonists."

DETERMINATION OF CITROVORUM FACTOR

Microbiological Assay. The microbiological assay for the citrovorum factor is carried out with Leuconostoc citrotorum (ATCC §8081) following the procedures proposed by Sauberlich 314

CITROLORUM FACTOR (LEUCOVORIN, FOLINIC ACID) STANDARD SOLUTION

⁽a) Preparation A water solution containing 100 pg DL-5-formy1-5, 6, 7, 8-tetra-

Sauberlich and Baumann J. Biol Chem., 176, 165 (1948).
 Flynn, et al J. Am Chem Soc., 73, 1979 (1951), Roth, et al J. Am Chem Soc.

<sup>74, 3247 (1952)

244</sup> Sauberheh and Baumann: J. Biol. Chem., 176, 165 (1918), Sauberheh: J. Biol. Chem.

^{181, 467 (1949)}

hydropteroylglutamic acid per ml is prepared. An aliquot of this standard solution is diluted in stages to 1 millimicrogram per ml (b) Assay Levels of Diluted Standard Solutions 0 1, 0 2, 0 3, 0 4, 0 5, 0 6, 0 7, 0 8,

09, 10 ml

BASAL MEDIUM FOR ASSAY OF CITROVORUM PACTOR WITH Leuconosloe calcotorum

Constituent	Amount per 500 ml double- strength medium	Constituent	Amount per 500 ml double- strength medium
	g		mg
Acid-hydrolyzed casein	10 0	Adenine sulfate	10 0
Sodium acetate	20 0	Guanine hydrochloride	10 0
Glucose	25 0	Uracil	10 0
Asparagine	0 10	Xanthine	10 0
Tryptophan	0 10	Thismine hydrochloride	0.5
Cystine	0 10	Riboflavin	0.5
NH-Cl	3 0	Calcium pantothenate	0.5
K,HPO.	0.6	Nicotinie acid	1 0
KH,PO,	0.6	Pyridoxine hydrochlo	1
	1	ride	1 0
Mg8O, 7H,O	0 2	Pyridoxamine hydro-	8
	1	ehloride	0 3
NaCl	0 01	Pyridoxal	0 3
FeSO, 7H,O	0 01	Biotin	0 001
MnSO, H1O	0 01	1	1

Combine ingredients adjust to pH 68 and make up to 500 ml with water

The preparation of the various constituents of the medium is as follows

VITAMIN-FREE, ACID-HYDROLYZEN CASEIN 245 Laboo "vitamin free" casein (300 g) is refluxed at least 12 hours with 3 liters 20 per cent HCl. It is evaporated under vacuum to a thick syrup, then diluted to 3 liters with distilled water. This concentration procedure is repeated twice The final concentrate is diluted to 1 liter The pH is adjusted to 25 with KOH. The mixture is stirred with 30 g of activated carbon (Darco G 60) for 15 minutes, then filtered with the aid of "Filter-cel" The pH is adjusted to 6 8 with KOH after the volume of solution is brought to 3 liters, and the charcoal treatment and filtration are repeated One inl of the solution contains the equivalent of 100 mg of original casein

ADENINE AND GUANINE These are dissolved together in a small amount of 1 0 N HCl with heat and diluted to 1 mg per ml

XANTHINE AND URACIL These are dissolved in dilute NH4OH and diluted to 1 mg per mI

TRYPTOPHAN Dissolve in a small amount of 1 0 N HCl and dilute to 10 mg per ml

ASPARAGINE One g of asparagine is dissolved in 100 ml of water

VITAMINS Ten mg thiamine hydrochloride, 20 mg pyridoxine hydrochloride, 6 mg pyridoxamine hydrochloride, 6 mg pyridoxal, 10 mg calcium pantothenate, 10 mg riboflavin, 20 mg nicotime acid and 0 02 mg biotin are dissolved in 100 ml distilled water and stored in the cold room under toluene. One ml of this solution is added per 100 ml of double-strength medium

¹⁴ Kitay McNutt and Snell J Bact 59, 727 (1950)

Preparation of sample solutions. Each sample for assay is blended with 150 ml. 0 05 M phosphate buffer (pl1 7.2) per g. of sample (dry basis). Caprylic alcohol is added to prevent foaming, and the mixture is autoclaved 15 minutes at 15 pounds. It is then cooled and incubated with desiccated chicken pancreas (Difco) using 20 mg. dry pancreas per g dry weight of sample under toluene for 24 hours at 37° C. After incubation the samples are autoclaved 5 minutes at 15 pounds, cooled, and filtered. The samples, together with the standard citrovorum factor dilutions, are pipetted into 12 × 100 mm test tubes and brought to a volume of 1 ml. with water, 1 ml. double-strength basal medium is added per tube. The tubes are capped with metal or glass caps and then are sterilized for 15 minutes at 15 pounds pressure. After cooling to room temperature, the tubes are inoculated as described below.

Preparation of the Inoculum. An Inoculum is prepared by growing the organism at 37° C, for 24 hours on the basal medium plus 10 mg Wilson's liver fraction L34 per tube. The cells are centrifuged, the supernatant liquud is discarded, and the cells are resuspended in 0.9 per cent saline. One drop of this suspension is added to 10 ml. sterile 0.9 per cent saline, and one drop of this cell suspension is added per tube. The assay tubes are incubated at 37° C for 20-24 hours. The growth response is measured by recording the optical density of the cultures with the photoelectric turbidimeter. From these data a curve can be plotted relating growth to citrovorum factor, and from this response curve the potency of the unknown samples can be calculated.

BIOTIN

A striking example of how the isolation of an active principle in pure form can lead to coordination of a vast amount of seemingly unrelated observations may be seen in the story of biotin Köğl and Tönnis²⁴ in 1936 reported the isolation from egg yolk of a yeast-growth factor (bios II) in the form of its methyl ester They assigned the name biotin to the free acid Recognition by West and Wilson²⁴⁸ of the probable identity of this factor with coenzyme R, a growth and respiratory stimulant for the legume nodule organism Rhizobium, was followed by the demonstration by György, Melville, Burk, and du Vigneaud²⁴⁹ of the identity of biotin with both coenzyme R and vitamin H The latter bad been previously reported to protect against a form of dermatitis induced by feeding raw egg white The identification of biotin also illustrates the contribution to the advance in nutritional science made possible by the application of microbiological methods. The structure of biotin was established by du Vigneaud and associates

In terms of the minimum protective dosage, biotin is one of the most potent physiological substances known, less than 0.03 μ g per day being sufficient for the rat. The role of biotin in nutrition is not clearly understood, but various investigations suggest that it may function as a coenzyme in certain enzymatic reactions associated with decarboxylation and carbon diovide fixation. The high concentration of biotin in embry-

other crude liver extracts may be used

²¹⁷ Kogl and Tonnis Z physiol Chem 212 43 (1936) 218 West and Wilson Science 39, 608 (1939) 219 Gyorgy Melville Burk and du Vigneaud Science 91 243 (1940)

onic and tumor tissue has attracted considerable interest in the possible clinical significance of this vitamin

Physiological Properties of Biotin. Biotin has been shown to be an essential nutrient for various lower organisms (including yeast, molds, bacteria and fungi) for the rat, chick, turkey, monkey, rabbit, dog, and guinea pig and for man Deficiencies of this vitamin are difficult to produce experimentally because it may be synthesized by intestinal flora in higher animals Deficiency symptoms may be induced, however, his feeding materials which either combine with the biotin to form non-absorbable complexes (such as avidin of egg white, see below) or by feeding sulfia drugs which interfere with bacterial synthesis of the vitamin in the intestines

In 1927 Boas discovered that the inclusion of large amounts of raw (but not of cooked) egg white in the diet of rats resulted in loss of hair, loss of weight, dermatitis and death. It was also found that certain foods could cure or prevent these symptoms. Raw egg white contains a distinctive protein, later designated as aridin capable of combining stoichiometrically with botin, thus preventing its absorption from the digestive tract or its utilization by yeast. Symptoms of a biotin deficiency may be induced even in man by feeding a sufficient amount of raw egg white Crystallized avidin (mol. wt. 70 000) has 15 000 times the biotin combining power of raw egg white. Though the biotin avidin complex is not alsorbed from the gastrontestinal tract, the compound is biologically active when administered parenterally

Biotin deficiency in animals is associated with the development of dermatitis, loss of fur disturbances of the nervous system, and death Biotin appears to be required for normal gestation and lactation in the rat and mouse as well as for normal development of the chick embry o A marked increase in the biotin content of the blood of sexually immature chicks was observed following the administration of estrogen, subsequent administration of progesterone caused no change in blood biotin but produced a rise in the avidin content of the oviduct These observations suggest a role of avidin and biotin in reproduction. The "spectacled eye" condition in rats (due to loss of hair around the eyes) may be associated with biotin deficiency In the chick the deficiency symptoms dermatitis and perosis are similar to those of pantothenic acid deficiency That vitamin interrelationships are involved in intestinal synthesis is illustrated by the observation that a pantothenic acid deficiency produced in rats on a diet containing succiny sulfathiazole could be relieved by administer ing folic acid and biotin 250

That botto may play a part in the synthesis of oleic acid is suggested by studies with certain lactobacillists. For example L arabinosus can multiply in the absence of botto provided oleate and aspartate are present Aspartic acid has a pronounced spaning action on the requirement of microorganisms for botto. If just sufficient botton is present to

Wr ght and Welch Science 97 426 (1943)
 Protter and Flychjern J Bud (Aem. 172 531 (1945)
 Williams and Fleger J Bud (Aem. 172 197 (1945)
 Williams and Fleger J Bud (Aem. 170 619 (194))
 Broquist and Snell J Bud (Aem. 188 431 (1951)

permit synthesis of oleie acid, aspartic acid can be replaced by oxalacetic acid. The microbial growth-stimulating effect of oleie acid is also manifested by certain detergents, e.g., Tween 80 (the polyoxyethylene derivative of sorbitan oleate). The significance of these observations in mammalian nutrition has not been established

The protective effect of liver or yeast concentrates against butter yellow (dimethylaminoazobenzene) hepatoma in rats has been found to be unrelated to their content of biotin. In fact pure biotin was reported to increase the incidence of tumors in susceptible mice ²⁵² Egg white affords some protection against tumors in rats, but this is independent of the biotin-avidin relationship ²⁵³ Addition of biotin to a diet containing butter yellow, but otherwise protective against tumor development, accelerated carcinogenesis; however, when the diet was conducive to tumor development, this effect was not observed ²⁵⁴

Most microorganisms require biotin in the form of the free acid, though the methyl ester is biologically active for yeast and partially so for Lactobacillus casei. The diaminocarboxyle acid obtained from biotin by the hydrolysis of the urea portion of the molecule possesses 10 per cent of the activity of the original vitamin Oxidation to the sulfoxide does not affect the activity of blotin, whereas conversion to the sulfoxide causes considerable inhibition of the growth stimulation of yeast Parenteral

milk It is also found in vegetables, grains, nuts, feeds, pollens, and molasses Biotin occurs naturally in combined forms, among which is a complex isolated from yeast and designated biocytin ** It is \$e\$-N-biotinyl-L-lysine Biocytin has the activity of biotin for L case; but not for L radbinosus Biotin exists in the free state in finits and grasses, partly bound in nuts, vegetables, and grains, and mostly in combined form in yeast and liver For assay with incroorganisms it may be liberated by autolysis, acid hydrolysis, or enzymatic digestion

Chemistry of Biotin.

cis-Hexahydro-2 oxo-I-H-thieno(3,4)imidazole-4-valeric acid

The structural formula shown is that of β biotin, isolated from liver or milk α -Biotin, from egg yolk, differs only in respect to the side chain which, instead of n-valeric acid, is —CH CH(CH₄)₂

COOH

The biotins are crystalline compounds soluble in water and alcohol and insoluble in chloroform, ether, and petroleum ether. The α - and β -biotins melt at 220° and 231° C, respectively They have specific rotations |a|25 of +51° and +91°, respectively, and an isoelectric point at pH 3-3 5 Biotin has an ultraviolet absorption maximum at 234 mm with a specific extinction coefficient of 42 5 It has a molecular weight of 244 3 The stability of biotin toward heat in the presence of acid or alkali has not yet been established It has been reported that brotin in natural materials resists autoclaving with strong mineral acids (4-6 N HCl for two hours at 120° C), a procedure used for the liberation of the vitamin in natural materials for microbiological assay Some authors, however, have reported a loss of biotin by this hydrolytic procedure. In natural materials the vitamin is destroyed by heating with strong alkali, and is even less stable to this treatment in pure solutions Both free and combined biotin are inactivated by oxidizing agents. Though esterification impairs the biological activity of biotin for microorganisms, acylation or alkylation do not affect it Synthetic \$\beta\$-biotin is produced commercially

In oxybiotin the sulfur atom is replaced by oxygen Oxybiotin possesses biological properties similar to biotin but in considerably less degree. It has about one-fourth the growth-promoting activity of botin for L helicticus and S cereusiae and half the activity for L arabinosis, it is one-third as active in curing botin deficiency in chicks and even less effective against egg-white injury in the rat

²²⁴ Wright et al J Am Chem Soc 72 1048 (1950)

Dethiobiotin, the natural preeursor of biotin, is believed to be derived in part from pimelic acid (COOH-(CH₂)s COOH). It can be made by reducing biotin with Rancy nickel, a reaction which leaves the furan ring intact.

Dethiobiotin is as active as biotin for yeast, E. coli, and various molds, but not for several species of lactobacilli, and has very little of the activity of biotin against egg-white injury in the rat.

DETERMINATION OF BIOTIN

Biotin may be determined microbiologically by measurement of its effect upon the growth of microorganisms. The method of Snell, Eakin, and Williams257 involves measurement of the growth stimulation of Saccharomyccs cerevisiac. The procedure is similar to that described on p. 1185. The extremely high physiological potency of biotin is illustrated by the fact that it stimulates yeast growth when present in a concentration as low as one part in 5 × 1011 parts of medium. Biotin may also be determined by its growth stimulation of other microorganisms including Lactobacillus arabinosus and Lactobacillus casei. A rat assay such as was used in studies of the anti-egg-wbite-injury factor (vitamin H) may also be employed, though it requires large groups of animals and is not as satisfactory as the microbiological determination. One mg of biotin methyl ester is equivalent to 27,000 units of vitamin H, the unit being the minimum daily dose required to cure the egg-white dermatosis in rats in 30 days. One great difficulty in the rat assay is that of producing biotin deficiencies, since rats obtain a considerable portion of their biotin from products of bacterial synthesis in their intestines. The assay for biotin using chicks is somewhat more satisfactory because they require greater amounts from dietary sources so that the deficiency is easily produced. Of the various assay methods, the microhiological procedure is generally preferred. See p. 1131.

VITAMIN B12 (COBALAMIN)

Vitamin B₁₅, the anti-pernicious-anemia factor of concentrated liver extracts and the major vitamin in sources of the "animal protein factor," was isolated in 1948 and found to he a cobalt coordination compound or "elielate." A second form, which was isolated in 1949, proved to be biolog-

ically interchangeable with the first I ater, these forms were identified as eyano (Bir) and hydroxo (Bir, = Bir, ***) forms of the organometable molecule containing cobalt. The known role of cobalt in the nutrition of ruminants could now be explained by the presence of this element in the molecule of these "cobalamins" which are synthesized by microorgainsms in the rumen. The identity of cobalamin with the "extrinsic factor" of pernicious anemia was demonstrated, and the production of cobalamin by industrial fermentation made a new source of the substance available in incheine and animal nutrition.

The term "vitamin B₁₂" refers chemically to cyanocobalamin but is commonly used to designate the cohalamin group with respect to their biological effect For example, the vitamin B₁₂ content of hier extract is commonly expressed in terms of a measurement of cyanocobalamin plus

hy droxocohalamin

Physiological and Clinical Aspects of Cobalamin. This vitamin is a growth factor for certain lattic acid bacteria and for the algal flagellate. Puglena gracilis The deficiency in young rats and chicks is seen when diets low in the vitamin are fed during the maternal period and is accentuated by raising the level of protein in such diets or by the addition of thyroid hormone to the diets fed to young animals. The deficiency in young rats is marked by a ligh mortality which is most prominent during the suckling period and is associated with leukopenia and uremia. The hatchability of the eggs of hens on deficient diets falls to low levels and there is slow growth and a high mortality in newly hatched chicks.

Pernicious anemia is the best-known example of the result of vitamin Bis deficiency In this disease, which is peculiar to the human species the patient by some degenerative process is deprived of an "intrinsic factor" which is present in normal gastric juice. This "intrinsic factor" is needed for the uptake of vitamin B12 ("extrinsie factor") from the digestive tract The disease is marked by a macrocytic anemia, by leukopenia, by megaloblastic changes in the bone marrow and usually by subacute com bined degeneration of the spinal cord. The signs and symptoms are reversed by administration of cobalamin As little as 1 or 2 µg daily will produce a therapeutic response when injected, as will 5 µg daily by mouth when given with intrinsic factor or a single dose of 3000 µg orally without intrinsic factor Pteroylglutamic acid will restore the blood and bone marrow pictures to normal in doses of a few milligrams daily when given either orally or by injection but this treatment fails to stem the progress of, or to alleviate the neural changes which respond only to cobalamin If, however these changes are not of recent origin, they are refractory to even large doses of cobalamin

Vitamin B₁₂ deficiency in pigs is marked by slow growth and is accompanied by nervousiess and irritability but not by macrocytic anemia. A wasting disease in ruminants known to be endemie in many areas was identified in 1935 as being due to a deficiency of morganic cobalt in the soil, with a consequent lack of this element in the forage. The disease in

sheep has been described as accompanied by listlessness, anemia, loss of appetite, and weakness progressing to a fatal termination. The disease can be prevented or arrested by cobalt salts when they are administered orally, but not when injected. Vitamin B_{12} is effective when either fed or injected, and it is thought that cobalt deficiency in the dict of ruminants leads to a failure in the fermentation process by which vitamin B_{12} is normally produced by the rumen microflora.

A source of the "labile methyl" group such as cooline, betaine, or dimethylthetin will enable rats to grow on a "labile-methyl-free" purified diet, containing homocystine and succinylsulfathiazole, without adding vitamin B₁₂ or pteroylglutamic acid. However, the rats will grow at a rate of 0 8 to 1.0 g. daily without a dietary source of "labile methyl" if both vitamin B₁₂ and pteroylglutamic acid are added to the diet. **so This relates these two substances to biological reactions involving precursors such as serine and glycine as sources of "formate," supplying a single-earhou fragment for the methylation of homocystine to form methionine. It was found by Shive** that the inhibitory effect of sulfanilamide on the growth of E. coli was overcome either by vitamin B₁₂ at a level of 0.3 µg, per liter of culture medium or by methionine at a level of 90 mg., but not by homocystine. A function of vitamin B₁₂ in the formation of methionine from homocystine was thus demonstrated.

Vitamin B₁₂ and pteroylglutamie acid both appear to "spare" the requirement of the chick for choline and to improve the utilization of

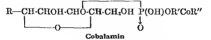
homocystine by the chick on methionine-deficient diets.

Vitamin B₁₂ appears to catalyze the formation of the purine and pyrimidine deoxyribosides which are present in deoxyrihonucleic acid. These deoxyribosides can replace vitamin B₁₂ in the growth of certain hacteria

including L. leichmannii.

The vitamin is formed by certain hacteria in the digestive tract and by soil and marine microorganisms. Its formation in the rumen doubtless accounts for its presence in beef liver. Vitamin B₁₁ is characteristically present in the internal organs and muscular tissues of animals and absent from the higher plants. The more important ductary sources of the vitamin are liver, bidney, lean meat, fish, clams, oysters, milk, and egg yolk.

Chemistry of the Cobalamins. The molecule consists of a large, highly stable cobalt complex (R'Co) esterified through phosphoric acid to a rubofuranoside group which in turn is linked through N to R, which is 5,6-dimethylbenzimidazole (as in vitamins B₁₇, B_{12b}, and B_{12c}) or adenine (as in "pseudo vitamin B₁₂"). An anion R", which may be CN



(in B₁₂), OH (in B_{12a}), NO (in B_{12a}), etc., and which is apparently not related to the biological activity of the substance, is attached by coordi-

²³⁹ Bennett · Science, 110, 589 (1949).

²⁴⁰ Shave: Ann. N. Y. Acad Sci , 52, 1212 (1950).

nate linkage to the cobalt atom Prolonged acid by droly six splits the phosphate linkages and al o liberates two p-1 amino-2 propanol groups from R' Catalytic hydrogenation of vitamin B. leads to the production of a brownish pigment which has an absorption band at 310 mu 241 Upon prolonged exposure to air the brown pigment becomes converted to hi droyocobalamin (vitamin Bira) which in turn may be changed to cyanocobalamin (vitamin B.) by treatment with evanide

The composition of vitamin B12 is approximately Car at H86 92 1401r PCo It forms dark red needle-haped crystals which in solution have absorption spectrum maxima at 278 361 and 500 mu The crystals of vitamin Bin are almost black and absorption spectrum maxima occur at 274 3.1, and a20 mg Both substances are readily soluble in water and insoluble in other They may be crystallized from acctone They are readily destroyed by alkalı Dilute acid converts B12 to B125 and strong acid causes by droly tic decomposition as described above The cobalamins are readily adsorbed on charcoal from which they may be eluted with appropriate solvents and they may be purified by chromatography oo siliere acid

An analog "pecudovitamin Bit ' was produced by anaerobic fermenta tion of an organism isolated from bovine rumen contents 262 Acid hydroly sis of this analog yielded adenine in place of the 56-dimethylbenzi midazole which is obtained from vitamin Bis the other hydrolytic prod ucts could not be distinguished from those yielded by vitamin Bi Pseudovitamin Biz was active for L leichmannis, E coli and Euglena gracilis hut was inactive in pernicious anemia and for chicks and rats Other microbiologically active derivatives of cobalamin mactive for animals have been detected in the feees of animals 244 These substances may be breakdown products of vitamin Bis

Determination of Vitamin B₁₂ This is usually carried out hy biological assay Among the organisms used are rats chicks Euglena gracilis I leichmannii and E coli mutant to 113-3 A correction for purine and pyrimidine deoxyribosides must be made if L leichmannii 15 used Chemical procedures have been described based on the addition of cyanide to cobalamin and the subsequent photolytic liberation of the evanide followed by its collection and determination

ml of sample in 25 ml of a freshly prepared bisulfite solution (13 fr g manchasic not essum phosphate and 10 g sodium metalisulfite dissolved in water and made to 1 liter) After cooling, dilute an abount of the clear supernatant with water so that the final test solution contains vitamin Bis activity approximately countries to 0.02 millimicrogram of evanocohalamin. The bisulfite aids in stabilizing naturally occurring analogs of the autamin. The amount of bisulfite is the assay tube should not as good 0.10 mg (or 0.02 mg per ml.)

STANDARD CYANOCOBALAMIN SOLUTION Prepare a stock solution by dissolving in 25 per rent alcohol an accurately weighed quantity of U.S.P. Cyanocoholaron Reference Standard, so that each ml contains 10 millimicrogram of evanocobolomic Oo the day it is to be used dilute 10 ml of the Stock Solution to 500 ml with vister Feeb ml contains 0.02 millimicrogram of expresshalamin

RIGHT MEDIUM STOCK SOLUTION

Acid-hydrolyzed Cascin Solution	25 ml
Cystine-Tryptophan Solution	25 ml
Asparagine Solution	5 ml
Adenine-Guanine-Uracil Solution	5 ml
Vanthine Solution	5 ml
Riboflavin-Thiamine-Biotin-Nicotinic Acid Solution	10 ml
p-Aminobenzoic Acid-Calcium Paatotheaate-Pyridoxiae-Pyridoxal	Pyridoxa-
mine-Folic Acid Solution	10 ml
Salt Solution A	5 ml
Salt Solution B	5 ml
Polysorhate 80 Solution	5 ml
Dextrose, Anbydrous	10 g
Sodium Acetate, Anhydrous	5 g
Ascorbie Acid	1 g

Dissolve the dextrose sodium acetate and ascorbic acid in the solutions, programely mixed add about 50 ml of water adjust to a pH of 6 0 with sodium hydroxide solution. and finally add water to make 250 ml

ACID-HYDROLYZED CASEIN SOLUTION Mix 100 g vitamin free casein with 500 ml dilute by drocbloric acid (1 in 2) and reflux the mixture for 8 to 12 hours Remove the bydrochloric acid from the mixture by distillation under reduced pressure until a thick paste remains. Redissolve the resulting paste in water, adjust the solution to a pH of 3 5 (±0 1) with sodium hydrovide solution, and add water to make 1000 ml Add 20 g activated charcoal, stir for 1 hour, and filter Repeat the treatment with activated charcoal, stir for 1 hour, and filter Repeat the treatment with activated charcoal Store under toluene in a refrigerator at a temperature not below 10° C. Filter the solution if a precipitate forms upon atorage

CYSTINE-TRYPTOPHAN SOLUTION Dissolve 400 mg each of 1-cystine and n.1tryptophan in 100 ml N HCl

ASPARAGINE SOLUTION DISSOIVE 20 g 1-asparagine in water to make 200 ml Store under toluene in a refrigerator

ADENINE-GUANINE-URACIL SOLUTION Dissolve 0 2 g each adenine sulfate, guanine hydrochloride, and uracil, with the aid of heat, in 10 ml 20 per cent hydrochloric acid.

cool, and add water to make 200 ml Store under toluene in a refrigerator AANTHINE SOLUTION Suspend 0 2 g vanthine in 30 to 40 ml water, heat to about

70° C , add 60 ml ammonia Test Soluting, and stir until the solid is dissolved Cool and add water to make 200 ml Store under toluene in a refrigerator

RIBOFLAVIN-TRIAMINE-BIOTIN-NICOTINIC ACID SOLUTION Prepare a solution in 0 02 N acctic acid, each ml to contain 25 pg riboflayin, 25 pg thiamine hydrochloride. 0.2 µg biotin, and 50 µg nicotinic acid Store, protected from light, under toluene in n refrigerator

p-Aminorphiole Acid-Calcius Panothenate-Pthiodenat-Pthiodenate-Pth

SALT SOLUTION A Dissolve 10 g monobasic potassium phosphate and 10 g dibasic potassium phosphate in water to make 200 mf Add 2 drops of hydrochloric acid, and

store under toluene

SAIT SOLUTION B Dissolve 4.0 g magnessum sulfate, 0.2 g sodium chloride, 0.2 g ferrous sulfate, and 0.2 g manganese sulfate in make 200 ml Add 2 drops of hydrochloric acid, and store under toliumen

POLYSORBATE 80 SOLUTION Dissolve 20 g polysorbate 80 in sufficient alcohol to make 200 ml Store in a refrigerator

TOWATO JUICE PREFARATION Centrifuge 1000 ml commercially canned tomato juice Suspend 5 to 10 g analytical filter and in the supernatant figured and filter under reduced pressure, through a layer of analytical filter and of sufficient thickness so that a clear, straw-colored filtrate is obtained 'store under tolurne in a refing rator

Cultume Medium Dissolve 0.75 g water-soluble yeast extract, 0.75 g peptone, 1 g anhydrous dextroee, and 0.2 g potsessum hiphosphate in 60 to 70 ml water Add 10 ml tomate pure preparation and 1 ml polysorbate 80 solution Adjust the solution to pH 63 with vodium hydrousde solution, and add water to make 100 ml Place 10-ml portions of the solution in test tubes, and plug with ection Sterlize the tubes and contents in an autoclase for 15 minutes at 121° to 123° C (exhaust line temperature)

SUSPENSION MEDIUM Dilute a measured volume of basal medium stock solution with an equal volume of water Place 10-ml portions of the diluted medium in test tubes Sterilize and cool as directed above for the culture medium.

Spock Cletter of the Leckbacellus leckmanns: To 100 ml culture medium add 10 to 1.5 g agar, and heat the matture, with sitting, on a steam bath, until the sgar dissolves Add approximately 10-ml, portions of the hot solution to test tukes, plug the tubes with cotton, stenlize for 15 minutes in an autoclave at 121° to 123° C (exhaust line temperature), and allow the tubes to cool in an upright position Prepare a stab culture from the pure culture of L techmanni, obtained from American Type Culture Collections** (No 7830) Incubate 6 to 24 hours at any constant temperature between 30° and 37° C Activate a newly obtained culture by making several daily transfers before use in an assay Maintain the culture in an active state by transfer at least three times weekly

INOCULUM Make a transfer of cells from the stock culture to a sterile tube of culture medium, and incubate 6 to 24 hours at a constant temperature between 30° and 37° C. Under aseptic conditions centrifuge the culture, decant the supernatant, and suspend the cells in 10 ml sterile suspension includes.

of the test solution of the material to be assayed. To each tune add 5.0 ml of basel medium stock solution and sufficient water to make 10 ml

After mixing, cover the tubes sultably to prevent bacterial contamination, and sterilize the tubes and contents in an autoclase for 5 minutes at 121° to 123° C (exhaust-line temperature) arranging to reach this temperature in not more than 10 minutes Cool as rapidly as practicable to avoid color formation resulting from overheating the medium. Take precautions to maintain uniformits of sterilizing conditions throughout the asset.

Asantically anocalate each tube fercent three tubes containing no Standard Cyanocobalamin Solution-uninoculated blank) with 1 drop of inoculum Incubate for 6 to 24 hours, until maximum turbidity is obtained, as demonstrated by a lack of significant change during a two hour nerlod in the tubes containing the highest level of Standard Cyanocobalamin Solution (0.1 millimleroteam) Maximum turbidity is predingrily obtained in 14 to 24 hours Read the turbidity of the tubes in a suitable instrument, at a specific wavelength that is ontimal for the Instrument used. This will lie between 540 and 660 m. In talling the instrument reading thoroughly mix the contents of each tube and transfer to ontical classware. Additate each tube or cuvette to obtain a uniform cuspension. A few seconds after adjustion a steady state is reached in which the daivangmeter needle remains constant for 30 seconds or more allowing sufficient time for an instrument reading. A little practice will establish the proper time interval With the moculated blank in the instrument, set the meter to read 100 per cent transmittancy, read the transmittance of the inoculated blank (the inoculated tubes to which no Stand ard Cyanocobalamin solution has been added). Disregard the results of an assay if contamination with a foreign organism is evident, or if the inoculated blank tubes give a reading of less than 90 per cent transmittancy (Evelyn or Lumetron), 80 per cent (Coleman), or 65 per cent (Reckman), thereby indicating interference due to vitamin Br activity in the basal medium stock solution or inoculum Then with the inoculated blank in the instrument set the meter to read 100 per cent transmittancy. Read the transmittancy of the tubes of the standard and sample series. Disregard the results of an assay if the transmittancy of the tubes containing the highest level of Standard Cyanocobalamin Solution (0.1 millimicrogram) is more than 65 per cent (Evelyn or Lumetron) or 50 per cent (Beckman or Coleman)

CALCULATION Prepare a standard concentration response curve by plotting the percent transmittancy readings for each level of the Standard Oyanocohalman Solution used against millimerograms of eyanocohalman contained in the respective tubes Draw the smooth curve which by visual inspection appears to fit best the plotted route.

From this standard curve determine by interpolation for each tube the amount of cyanocobalamin equivalent to the vitam n B activity of each ml of the test solution of the material to be assayed.

Since in microbial assays occasional inexplicable aberrant values are obtained in individual tubes inspect the series of values and set aside any which vary markedly from most of the series Strike a provisional average of the remaining values and set aside any of the latter which are less than 90 per cent and more than 110 per cent of the provisional average II less than 10 of the 15 original values remain the data are insufficient for calculating the potency if 10 or more values remain calculate the potency from the average of them. If the calculated potency of an assay is less than 75 per cent or more than 125 per cent of the 48 med potency, the result is out of the entitied range for greater accuracy a reassay at a level more closely approximating the true value is necessary.

Determination of Cobalamin (Pad-Plate Method of Williams and Coworkers):24 Principle. Cobalamin is determined by measurement of growth of E. cols mutant The diameter of the growth zones on agar plate is measured and evaluated on a standard curve.

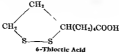
Distribution of Cobalamins. Good food sources include liver, kidney, lean meat, fish, milk, and oysters The higher plants contain little or no vitamin B12, but it is produced by many microorganisms, including molds of the Streptomyces group which are used for its industrial production. Ruminating animals appear to obtain the vitamin from fermentations which are carried out by the microflora of their digestive tract; this accounts for the presence of cobalamins in heef liver and also for the fact that in ruminants a lack of cohalt leads to a wasting disease caused by a deficiency of vitamin B12.

THIOCTIC ACID (PROTOGEN, LIPOIC ACID)

This coenzymatic factor has not been shown to be essential in the diet of animals and should not, strictly apeaking, be classified as a vitamin. However, thioctic acid has biochemical functions which relate it closely to the B vitamins.

The protozoon Tetrahymena geleii was shown by Dewey and Kidder256 to need an unidentified factor present in liver and other natural materials. This factor was shown to contain a component which differed from the known vitamins and was named "protogen" Two chemically distinct forms were shown to exist.247 In other investigations extracts prepared from yeast were found to be interchangeable with acetate in promoting the growth of L. caser248 and to be needed for the oxidation of pyruvate by resting cells of a strain of S. fecalis,269

The crystallization and study of one form of the acctate-replacing factor, which was renamed "α-lipote acid," showed the presence of sulfur²⁷⁰ and a similar compound obtained from liver was also found²⁷¹ to contain sulfur The synthesis of DI-6,8-dithrocetanoic acid (abbreviated to "6-thioctic acid") was accomplished following studies of the structure of the natural compound 272 The biological activity of 6-throctic



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 Dewey Proc Soc Exptl Biol Med 46, 482 (1941), Biol Bull., 87, 107 (1944).

Kidder and Dewey Biol Bull, 87, 121 (1944)

101 Stokstad, Hoffman, Regan, Fordham, and Jukes Arch Biochem, 20, 75 (1949).

¹⁴⁴ Guirard, Snell, and Williams Arch Biochem, 9, 381 (1946).

²⁴⁵ O'Kane and Gunsalus J Bact . 54, 20 (1947) m Reed, et al. Science 114, 93 (1951), J. Biol. Chem., 192, 851, 859 (1951)
11 Patterson, et al. J. Am. Chem. Soc., 73, 5919 (1951)

¹¹² Bullock, Brockman, Patterson, Pierce, and Stokstad J Am Chem Soc., 74, 3455 (1952).

acid is high, 0.4 parts per billion of purified culture medium is sufficient. to produce half-maximum growth of T gelen It was suggested that the compound may be present in hiological systems in acid amide linkage with thismine pyrophosphate 273 Oxidation and reduction of 6 thiortie acid give rise respectively to an oxide (protogen B or 8-lippor acid) and a dithiol

Physiological Aspects of Thioctic Acid. The interchangeability of protogen with acetate in the nutrition of S. fecales and the role of protogen in the oxidation of pyrilyate were early observations which indicated the coenzymatic functions of the substance and its probable relation to cocarbo vlase (diphosphothiamine) Approximately 6 moles of protogen and 1 mole of diphosphothiamine were found to be present in α-ketoglutarie acid o'idase, *71 and the presence of protogen in pyruvic oxidase was also noted, *73 suggesting a possibility that steps of the following type might occur in the oxidation of keto acids *76

R CO COOH + thiamine
$$\rightarrow$$
 R COH thiamine + CO₂

CH₂

CHR' \rightarrow R CO S(CH₂)₂CH(SH)R + thiamine

HS(CoA)

HS(CH₂)₂CH(SH)R + RCO S(CoA)

DPN+ |R|S

Oxidation of (B), the dihydrotbioctic acid form of the coenzyme, restores the latter to its original condition

Chemistry of Thioctic Acid. 6 Thioctic acid is a light-vellow crystalline solid, M P 60°-61° C (Fig 283) It is insoluble in water but readily dissolves in alcohol, acetone, ether, and many other organic solvents The sodium salt is freely soluble in water Thioctic acid is comparatively stable, but solutions tend to become oxidized by air to thiosulfinyloctanoic acid (oxythioctic acid, protogen B, or \$ lipoic acid) The ultraviolet absorption spectrum shows a peak at 332 m_µ (molar extinction coefficient ±160) Calvin and Massini 77 have suggested that light shifts the equi librium of the thioctic-acid containing coenzyme toward the reduced (dithiol) form, thus rendering it meanable of oxidatively decarboxy lating pyruvic acid (newly formed from CO2) to give rise to acetyl coenzyme A required to bring this carbon into the tricarboxilic acid excle. It was

²⁷¹ Reed and DeBusk J Biol Chem 199 881 (1952)

^{*} Sanadı Littlefield and Bock J Biol Chem 197 851 (1952)

<sup>177 851
171</sup> Schweet and Cheslock J Biol Chem 197 749 (1952)
171 Reed and DeBush J Am Chem. Soc 75 1961 (1953)
177 Calvin and Massini Experientia 8 445 (1952)

further suggested "* that a bradeal formed by dissociation of the disulfide bond in a strained five-membered disulfide containing ring, such as in thioctic acid, is the form in which light energy travels from chlorophyll to the chemical reactions concerned with the fivation of carbon dioxide in photosynthesis (see also p. 58)



T- - 900 Turnman to

ADENING AND GUANINE These are dissolved together in a small amount of 1.0 N HCl with heat and diluted to 1 mg, per ml

Unactt. Dissolve in dilute MI-OH and dilute to 1 mg per ml

TRYPTOPHAN Dissolve in a small amount of 10 N HCl and dilute to 10 mg per ml
ASPARACINE Dissolve I g apparague in 100 ml water

ASPARCIAE Dissolve 100 mg 1 mostol, 100 mg choline chloride, 10 mg calcium pantotherate, 10 mg thymine hydrochloride, 10 mg nicotinamide, 10 mg riboflavia.

BASAL MEDICAL FOR ASSAY OF THEOCHE ACID WITH Corunehacterium homes

Constituent	Amount per 200 WI Double- Strength Medium	Constituent	Amount per 200 Ml Double- Strength Medium
Acid by droly zed casein	5 g	ı Inositol	5 mg
Glucose	5 g	Choline chloride	5 "
Sodium acetate	2 5 g	Calcium pantotlienate	05 "
Speakman salt solution			
A	5 ml	Thiamine	05 "
Speakman *alt solution	ł	1 1	
В	5 ml	Nicotinamide	05"
Asparagine	50 mg	Riboffavin	05'
pt-try ptophan	100 mg	Pyridovine	05 "
L-cy stine	100 mg	p-Aminobenzoic seid	05 "
Adenine sulfate	10 mg	Pteroylglutamic acid	05 "
Guanine hydrochloride Uracil	10 mg 10 mg	Biotin	05"

Combine ingredients adjust to pH 68 and make up to 500 ml with water

10 mg pyrdovne hydrochloride, 10 mg p-aminobenziocacid, 10 mg pteroylglutamic acid, and 10 mg biotin in 100 ml distilled water, and store in the cold room under toluene Add I ml of this solution per 100 ml of double-strength medium

Preporotion of Somple Solutions 1f the thloctic acid in the test sample is in a "hound form," the sample should be autoclaved at 15 pounds pressure for 2 hours with 2 volumes of 8 N sulfuric acid, cooled, neutralized, and filtered to give a clear filtrate for assay. The assay samples, together with the standard dilutions of thoetic acid, are piperted into 12 × 100 mm test tubes and brought to a volume of 1 ml with water, 1 ml double-strength hasal medium is added per tube, the tubes are capped with metal or glass caps, and then are sterilized for 15 minutes at 15 pounds pressure. After cooling to room temperature the tubes are inoculated as described below

Preporotion of the Inoculum: An inoculum is prepared by growing the organism at 25° C for 48 hours on the basal medium plus 10 mg Wilson's liver fraction L^{ml} per tube. The cells are centrifuged, resuspended in an equal volume of saline, and 1 drop is used per 2-ml assay tube. The assay tubes are incubated at 25° C for 72 88 hours. The growth response is measured by recording the optical density of the cultures with the photoelectric turbidimeter. From these data is curve can be plotted relating growth to thioctic acid concentration, and from this response curve the potency of the unknown samples can readily be calculated.

²²¹ Other crude liver extracts may be used

Distribution of Thioctic Acid. The substance is widely distributed in biological materials Liver, pancreas, yeast, green leaves, and soybean meal have been reported to be good sources The throctic acid content of the chick embryo increases rapidly during the final days of incubation, presumably indicating synthesis by the tissues

PARA-AMINOBENZOIC ACID

The earliest indications of the nutritional importance of p-aminobenzoic acid were observations that the compound counteracted the hacteriostatic effect of sulfanilamide 282 Its importance in the nutrition of microorganisms and of higher animals was established shortly thereafter, as well as

its presence in certain natural materials particularly yeast 282

Physiological and Clinical Aspects of p-Aminobenzoic Acid. p-Aminobenzoic acid is the "bridge" between pteroic and glutamic acids in the structure of folic acid Although p-aminobenzoic acid (PABA) is frequently classed among the vitamins there is little to justify the belief that it plays a direct role in human nutrition Evidence points rather to its essential role in microhial nutrition and hence to its stimulating effect on vitamin synthesis by intestinal microflora Direct evidence of such bacterial synthesis is seen in the observation that excretion of p-aminobenzoic acid greatly exceeded dietary intake (See p 812)

An excess of n aminobenzoic acid inhibits the bacteriostatic effect of sulfonamides possibly because of structural similarities (see Chapter 36) This antagonism obeys the mass action law, thus indicating that the inhibition is competitive. The antisulfonamide effect of the vitamin has been noted in vitro 154 as well as in vito in mice infected with Streptococcus hemolyticus Sulfonamide resistant strains of Staphylococcus have been developed in which the resistance is proportional to the ability of the microorganism to synthesize p aminobenzoic acid p-Aminobenzoic acid interferes with the malaricidal action of sulfanilamide drugs but not with that of quinine and atropine, which probably attack the microorganism through a different channel The use of p aminobenzoic acid derivatives as local anesthetics and their possible incompatibility with sulfonamides ad ministered subsequently as bacteriostatic agents has been considered Observations relating to the hehavior of p-aminobenzoic acid under physiological conditions however indicate that p-aminobenzoic acid therapy probably does not interfere with subsequent administration of sulfonamides Furthermore it has been found that small amounts of p-aminobenzoic acid often potentiate sulfonamides

maintenance of hair color in the black rat. 288 The injection of hydroquinone in cats or mice results in graving of the fur, which can be prevented or cured by n-aminobenzoic acid. The use of PABA in the treatment of nutritional achromotrichia in man, reported to be successful in a few isolated cases has been refuted on the basis of more carefully controlled studies.259 Anti-gray-hair properties have also been demonstrated for pantothenie acid, biotin, and folic acid, Stimulation of the growth of microorganisms by the administration of one of the vitamin B complex may cause an increased synthesis or utilization of another member by the intestinal flora. This makes it difficult to ascribe particular physiclogical effects to specific B vitamins when administered orally. Thus the action of n-aminobenzoic acid in the prevention of achromotrichia has been ascribed by some observers to alterations in the intestinal flora promoting the synthesis of pteroviglutamic acid or of other factors coneerned with melanin formation.

In some species, p-aminobenzoie acid has been found to increase the physiological potency of insulin and penjeillin. It may also play a role in the metabolism of hormones—for example, by inhibiting the production of thyroid hormones. In addition, the rate of enzymatic inactivation of stilbestrol by mushroom tyrosinase, as well as the oxidative destruction

of epinephrine, are inhibited by n-aminobenzoic acid.

p-Aminobenzoic acid absorbs ultraviolet radiations (maximum 297.5 mu) in the range which produces sunburn and suntan in human slin

p-Aminobenzoic acid participates in certain detoxication reactions. In rats PABA detoxifies high doses of pentavalent and trivalent arsenical drugs used in the treatment of various forms of syphilis, without interfering with their bacteriostatic potency. It has also been claimed to detoxify antimony compounds used in tropical diseases.

Large doses of p-aminobenzoic acid are toxic to dogs and mice. The oral administration of more than I g. per kg. body weight in dogs is fatal. On the other hand, the administration of 1.4 g. per kg, body weight to rats is nontoxic. Moderate doses are acetylated by man and excreted in

Distribution of p-Aminobenzoic Acid.296 p-Aminobenzoic acid is present in most tissues. It occurs in nature in both the free and the combined form. Relatively high concentrations (several micrograms per gram) are found in yeast, liver, rice bran, rice polishings, and whole wheat, In the last it is present mostly in the germ. Milk contains about 0.1 mg. per liter.

Chemistry of p-Aminobenzoic Acid, p-Aminobenzoic acid was synthesized by Fischer291 as early as 1863, by reduction of p-nitrobenzoic acid with ammonium sulfide. It crystallizes in colorless needles which

²¹⁸ Martin and Anebacher: J. Biol. Chem., 138, 441 (1941).
²¹⁹ Brandalcone, Main, and Steele. Proc. Soc Expll. Biol. Med., 53, 47 (1943); Eller and Diaz. N.Y. State J. Med., 43, 1331 (1943)

²⁹⁰ For quantitative data see Landy and Dicken J. Biol. Chem., 146, 109 (1942); Mitchell, Isbell, and Thompson told , 147, 485 (1943) and 148, 281 (1943). mi Fischer. Ann , 127, 142 (1863).

melt at 186° C. It is soluble in water to the extent of 0.5 per cent at room temperature, but is freely soluble in alcohol or boiling water



2 Aminobenzoic acid

DETERMINATION OF p-AMINOBENZOIC ACID

No satisfactory chemical method is available for the determination of p-aminobenzoue acid. It reacts with p-dimethylaminobenzaldehyde in glacial acetic acid to produce a compound having a yellow color. However, this reaction, as well as other coupling reactions, is nonspecific and is given by isomers of p-aminobenzoue acid as well as by aniline and its derivatives, and by sulfonamides in the analysis of biological materials the bound vitamin must first be freed by strong acid or alkaline by drolysis.

p-Aminobenzoic acid is required for the growth of several microorganisms, including Brucella abortus, Streptococcus hemolyticus, and Clostradium acetobutylicum. Microbiological methods for the determina tion of PABA are hased upon the growth stimulation of Acetobacter suboxydans** and of a p-aminobenzoicless mutant of Neurospora crasar Determinations may also be made employing organisms which do not require an external supply of the vitamin by measuring the degree of inhibition of the hacteriostatic effect of sulfa drugs resulting from the addition of p-aminobenzoic acid

CHOLINE

Choine has occupied a prominent place in biochemical literature because of its relationship both to the phospholipides and to its acet'd seter Acet'd lednine, first studied as a synthetic product, then found in plant and animal tissues, plays an important role in the bumoral transmission of parasympathetic and other nerve impulses to effector organs. More recently, choline has achieved added significance hecause of its role in the process of transmethylation (see below and p. 1029)

Choine, prohably of pho-pholipide origin, was isolated independently by numcrous investigators and has been variously designated sinkalin hillneume fagin, amaintin and neume the latter term being now reserved for the unsaturated hase trimethylvinylammonium hydroxide

Because of its biochemical function and its distribution in foods choline is usually considered along with the vitamins of the B group However it not only lacks the specifieity characteristic of the vitamins but is actually a structural component of fat and nerve tissue. Moreover it is not known to participate as a cofactor in an enzyme system. Hence tholine is not strictly speaking a vitamin Part of the syndrome of

¹¹ Cheldelin and Bennett J Biol Chem 161 751 (1945) 11 Agaiwala and Peterson Arch Biochem 27 304 (19.0)

choline deficiency in animals can be cured by the administration of other methyl group donors, (e.g. methionine, betaine) or of vitamin B_{12} which

plays a role in methyl group transfer.

Physiological Properties of Choline. Choline is essential for normal growth of the rat, chick, and dog, and for lactation in the rat. Dictary deficiency has been demonstrated to be responsible for paralysis in the hind legs of nursing rats, inhibition of egg production in hens, slipped tendon (perosis) in chicks and young turkeys, and fatty liver in rats, dogs, rabbits, and pigs. The last-mentioned symptom is not observed in guinea pigs, whose low requirement for choline is attributed to a lack of hepatic choline oxidase. Choline deficiency in young rats produces an acute hemorrhagic lesion of the kidneys. Even after short exposure to this condition and subsequent restoration to an adequate diet, such rats have heen reported later in life to develop hypertension with eardiac enlargement and persistent renal damage.²³⁴

Choline performs several physiological functions. It enters into the molecular structure of phospholipides and actylcholine, and supplies lahile methyl groups for transmethylation reactions. Phospholipides are concerned with the mobilization of fat in the body. In the absence of choline, neutral fat, and to some extent cholesterol esters, accumulate in the liver. Choline is also lipotropic in that it prevents fatty livers in depancreatized dogs. However, this condition, when induced in rats hy

high-cholesterol feeding, does not respond to choline feeding.

Certain specific compounds with lahile methyl groups, or otherwise related in structure, can replace choline in some of its biological functions Methionine mobilizes liver fat in a manner similar to choline, and both the p- and 1-forms are equally effective. The mechanism probably involves the transfer of labile methyl groups from methionine to ethanolamine with the formation of choline. In support of this hypothesis is the observation that choline containing deuterium may be isolated from the animal organism after feeding methionine whose labile methyl group contains deuterium. Betaine and other linetropic factors may also supply methyl groups to ethanolamine. Conversely, methionine, an essential amino acid, can be replaced by homocystine when choline, betaine, or dimethylethylammonium ehloride is fed. The methyl-diethyl and the triethyl homologs do not support growth, but are strong lipotropic agents, and prevent the occurrence of hemorrhagic kidneys. Arsenocholine and sulfocholine likewise have lipotropic action and prevent renal hemorrhage, but neither of these compounds can methylate homocysteine. There is no diminution of activity when phosphorus is substituted for the nitrogen of choline. The hydroxyl group must be free, however, since ethers are inactive. The methyl groups of creatine. S-methyl cysteine, or of the hetaines from threonine, serine, or allothreonine, are not available for transmethylation.

Choline serves as a methylating agent in the physiological process: guanidoacetic acid → creatinine Here, too, methionine and betaine can replace choline.

²⁷⁴ Hartcroft and Best; Best Med J , 1, 423 (1949).

Cholme is involved in the transmission of nerve impulses in the form of its acetyl ester, acetylcholme a powerful agent which lowers the blood pressure. The organism maintains a delinate balance between acetylcholme and cholme (the latter having only slight vasodilator effect) by means of specific enzyme systems which can acetylate to acetylcholme or hydrolyze to free cholme when necessary.

Storage and Synthesis of Choline. Choline is present in the animal organism chiefly in the form of the phosphatides, i.e., the lecithins and splingomyelins. The fatty matter of brain kidnes, and liver is distinguished from that of adipose tissue by the fact that it consists largely of phosphatides. Lecithin is the glycryl ester of a pair of fatty acids and a substituted phosphoric acid group attached to a choline radical Sphingomyelin contains a fatty acid splingosine, and a choline-phosphoric acid group. In bect liver, only 2 per cent of the total choline is present in the free form

As discussed above choline may be synthesized in riso from cthanol amine and a methyl donor, methionine or betaine Rats on a diet containing ample methionine, for example need no dietary supply of choline

Distribution of Choline "The concentration of choline in animal tissues is proportional to their phospholipide constituents. Thus egg yolk liver, kidneys, brain, heart and nervous tissue are rich sources Good sources of choline are muscle tissue green, leafy and leguminous vegetables seed oil meals, and grain germs. In corn meal, 75 per cent of the choline is concentrated in the germ while 50 per cent of the choline in wheat is lost in the preparation of white flour. In general, seed meals and seeds are considerably better sources than the ecreal grains. Butter, lard, and refined vegetable oils are almost devoid of choline. Vixed diets of man have been shown to provide 0 to 10 g of choline per day, an order of magnitude more nearly resembling that of individual amino acids than that of vitamins.

Chemistry of Choline Choline was first isolated by Strecker in 1849 and synthesized by Wurtz in 1867. It possesses the following structure



The free base is a colorless crustalline extremely hygroscopic compound it is a stronger base than ammonia and easily forms salts with acids. Pure choline when heated decomposes to trimethylamine and ethylene gly col. Aqueous solutions containing less than 4 per cent choline may be heated without decomposition but losses occur in more concentrated media. Choline is more stable to heat in acid than in neutral or alkaline solution. Choline dihydrogen citrate, choline chloride, and carbamyl choline chloride are used medicinally.

Choline is readily soluble in water, methyl alcohol, ethyl alcohol, and formaldehyde; slightly soluble in dry amyl alcohol, dry acetone, and chloroform; and insoluble in dry ether, petroleum ether, benzene, toluene, earbon bisulfide, and carbon tetrachloride The salts are soluble in water and alcohol, forming aqueous solutions which are nearly neutral.

Choline may be precipitated from aqueous solution by potassium triiodide, phosphotungstie acid, phosphomolybdic acid, or reinecke salt. These precipitants, especially the last, have been employed in quantitative methods for its determination. Choline chloride also forms characteristic double salts with the chlorides of platinum, gold, and mercury.

DETERMINATION OF CHOLINE

Microbiological Assay (Method of Horowitz and Beadle):*** Principle. Choline may be measured by the growth stimulation of a mutant strain of Neurospora crassa designated as choineless, produced artificially by exposure to ultrau nolet radiation. Methodness, which can replace choline for the microorganism, is removed by adsorption of the extract on "Permutit" followed by elution with sodium chloride solution. The microbiological procedure for the determination of choline is far more sensitive and possibly more specific than the chemical procedure **The latter involves precipitation and isolation of the reineckate, followed by colorimetric measurement of the red pigment at \$20 mg in acctone solution.

Procedure, Heat 100 mg, of sample in an autoclave for 2 hours at 15 pounds pressure in 10 ml. 3 per cent sulfuric acid, or reflux for 7 hours, Neutralize to congo red with saturated barlum hydroxide solution. Centrifuge and filter the supernatant through a Whatman No, 50 paper. Add 3 ml. distilled water to the precipitate and bring to a boil with stirring. Gool, centrifuge, and add the washing to the previous supernatant. Neutralize the solution to litmus with 1 N sodium hydroxide solution and dilute with distilled water to a concentration of approximately 15 µg, per ml. Pass 5 ml. of the neutralized extract*31 through a column 100 mm. long and 5 mm. wide (internal diameter) containing 1 g. of Permutit. Wash the column with 5 ml. of 0.3 per cent sodium chloride and discard the filtrate and washings. Elute the choline with 10 ml. of 5 per cent sodium chloride, *30 per cent sodium chloride, *30 per cent sodium chloride.

Prepare a basal medium having the following composition in g. per liter: ammonium tartrate 5, ammonium nitrate 1, monobasic potassium phosphate 1, magnesium sulfate-7H₂0 0.5, sodium chloride 0.1, calcium chloride 0.1, calcium chloride 0.1, sucrose 20, biotin 5 × 10⁻¹. In addition, add the following trace elements as saits in mg. per liter: boron 0.01, molybdenum 0.02, iron 0.2, copper 0.1, manganese 0.02, and zinc 2.0.

Maintain stock cultures of the cholineless mutant No. 34486 of Neurosporo crassa on agar siants composed of the basal medium plus 1.5 per cent agar, 0.2 per cent Difco yeast extract, 0.2 per cent malt extract, and 1 pg. per ml. of choline. For inoculum, prepare a spore suspension in a few ml. of distilled water.

Pipet 0.5, 1.0, and 2.0 ml. aliquots of the sodium chioride cluate in duplicate into 250-ml. Erlenmeyer flasks, and dilute each to 25 ml. with basal

²³⁴ Horowitz and Beadle: J. Biol. Chem , 150, 325 (1943).

¹¹⁷ U S Pharmacopoeta XIII

m If the solution is known to contain less than 3 µg. of choline per ml , pass 10 ml through the column.

²³¹ If more filtrate is required in case the approximate choline content is unknown, adsorb

medium The final concentration of choline should lie between 2 0 and 20 pg per 25 ml At the same time set up a series of standards having 0 2 4 8 12 16 20 25 and 30 µg of choline per flask Autoclave the solutions at 15 pounds pressure for 10 minutes cool and add 1 drop of inoculum to each incubate at 25° C for 3 days At the end of this period collect the mold pads on tared fritted glass filters of medlum porosity and wash with distilled water Dry in an oven at 90° C and determine the dry weight

CALCULATION Prepare a reference curve from the data obtained with the standard ser es plotting µg of chol ne as the absc ssa and ve git of dry mold as the ordinate

From the scurve read off the et oline concentration C per mt of cluate Then

$$C \times \frac{10}{\Lambda} \times \frac{V}{0.1} = \mu g$$
 choline per g of sample

where I is the volume to hich the original extract as 11 ited and A is the volume of the al quot which vas passed through the Permitt column

INOSITOL

Though mositol was long known to be a constituent of heart muscle and of many plants (in the form of phytin the calcium magnesium salt of mositol heraphosphate) its functional similarity to the vitamins was not recognized until recently Eastcott200 isolated from yeast the factor previously known as bios I and demonstrated its identity with inositol The biologically active compound is optically mactive (designated mositol or meso mositol) as distinguished from its biologically mert stereosomers. In addition to its role as a yeast growth factor inositol has been identified with the mouse alopecia factor of Woolley and tho rat anti spectacled eye factor of Paveck and Baum 202

Physiological Properties of Inositol Though the metabolism or mode of action of inositol has not yet been elucidated its importance in the nutrition of several species has been demonstrated. An external source is required by some yeasts and fungi but not by all Young mice on a deficient diet suffer an inhibition of growth and loss of hair both of which are corrected by supplementation with mositol Growth of a transplanted mouse tumor has been inhibited by inositol It is also essential for proper growth in the chick and rat. The spectacled eye condition in rats due to loss of hair about the eyes is a result of dietary deficiency The vitamin also has lipotropic properties for this species but more for the cholesterol than the fat type of fatty livers Inositol stimulates gastrointestinal peristalsis in dogs Though the vitamin must be present in the free state to be available to yeast it is readily utilized by the mouse in natural bound forms such as phytin and soybean cephalin as well as synthetic methyl inositol and inositol hexagetate

The metabolism of mositol is related to that of other vitamins. It has been suggested that pantothenie acid regulates absorption of inositol from the intestine for a diet containing inositol produces symptoms of

¹⁰⁰ Eastcott J I hys Chem 32 1091 (1928) ¹⁰¹ Woolley J Biol Chem 136 113 (1940) ¹⁰² Laycek and Ba m Seance 92 384 (1940)

inositol deficiency if pantothenie acid is lacking. A relationship to p-aminobenzoic acid has also been established. In one series of experiments, black rats showed little signs of deficiency when both inositol and p-aminobenzoic acid were either absent or present, but exhibited typical symptoms when either nutrient alone was eliminated from the diet.

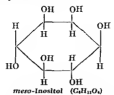
Because of its similarity in structure both to the carbohydrates and to the aromatics, inositol has been designated as a possible physiological

link between these two classes of compounds.

Storage and Synthesis of Inositol. The wide distribution and variety of forms of inositol in animal and plant tissues indicate that it is present for some physiological purpose rather than merely for storage of phosphoric acid or hydroxyl groups. It has been reported that inositol may be synthesized by intestinal flora, a process stimulated by the presence of pantothenic acid.

Distribution of Inositol. *e** In animal tissues, bigh concentrations are found in kidney, heart, spleen, thyroid, and testes. Inositol is found in the free form in muscle and other tissues; hence the name muscle sugar. In liver and heart muscle it is found combined, probably with a protein. Excellent dietary sources are cereal brans and seeds (in the form of phytin) and fruits, particularly of the citrus variety. A compound of inositol has been found in the phosphatide fraction of the soybean.

Chemistry of Inosifol. meso-Inositol is one of the eight cis-trans isomers of bevabydrocyclohevane, one of which occurs in optically active forms, making nine in all. It is a sweet-trasting crystalline compound and is isomeric in molecular formula with the bevose sugars. It crystallizes as the anhydride (M.P. 225°-226°) from water at temperatures above 50° C. and from anhydrous solvents. Below 50°, the dihydrate (M.P. 215°-216°) appears The molecule is stable to strong acid and alkaline hydrolysis. It is very soluble in water but insoluble in absolute alcobol or ether. meso-Inositol is optically inactive.



Determination of Inositol. The microbiological method for determination of vitamin B₆, depending on the growth stimulation of Saccharomyces cerevisiae (p. 1185) or carisbergensis³⁰⁴ may be used for the determination of inositol by simple modification of the basal medium to exclude juositol and include an excess of pyridovine. High specificity is

³³ For quantitative data see Woolley J Biol. Chem., 186, 453 (1941) and 147, 531 (1943); Scalock and Livermore J. Nutrition, 25, 265 (1943); Mollgaard, Lorenzen, Hansen, and Christensen: Biochem. J., 40, 589 (1948).
³⁴ Athun, et al.: Ind Eng. Chem., Anal. Ed., 15, 141 (1943).

medium. The final concentration of choline should lie between 2.0 and 20 ed ner 25 ml At the same time set up a series of standards having 0, 2, 4, 8. 12 16, 20, 25, and 30 up of choline per flast. Autoclave the solutions at 15 nounds pressure for 10 minutes, cool, and add 1 drop of inoculum to each Incubate at 25° C for 3 days At the end of this period, collect the mold pads on tared, fritted glass filters of medium porosity, and wash with distilled water Dry in an oven at 90° C and determine the dry weight

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$$C \times \frac{10}{A} \times \frac{V}{0.1} = \mu g$$
 choline per g of sample

where L is the volume to which the original extract was diluted, and A is the volume of the about which was passed through the 'Permutet column

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⁸⁴ Easter tt. J. Phys. Chem. 32, 1894 (1978) 81 Woolley. J. Biol. Chem. 33, 113 (1940)

^{#1 |} avcek and Daum Seance 92 394 (1940)

doses of ascorbic acid showed diminished skin hypersensitivity to arsphenamine 214 The protective effect of ascorbie acid against arsenical toxicity has been reported to be greatly augmented by the simultaneous administration of the flavonoid derivative, hesperidin methyl chalcone 215

An increased requirement for vitamin C in man has been reported in Hodgkin's disease protracted fevers active rheumatic heart disease and tuberculosis In rats the synthesis and excretion of ascorbic acid is stimulated by feeding certain cyclic Letones related to the terpenes and certain simple aliphatic ketones 216 This increased synthesis may be related to greater requirements for detoxification purposes. The beneficial effects of vitamin C in the detoxification of other unrelated poisons have been reported, eg, lead and arsenic compounds, benzene, and the toxins of several pathogenic microorganisms

Under normal conditions a dietary supply of ascorbic acid is required only by man other primates, guinea pigs, and some microorganisms However, the vitamin is essential for the normal development of most species, and is produced endogenously, thus resembling a bormone Interference with the normal synthesis of the vitamin in the bovine species results in the impairment of the reproductive functions of both the male and female Dietary deficiency of vitamin A results in decreased synthesis of ascorbic acid in the rat, though this relationship with vitamin A has not been observed in mature hens or bulls. In the latter species tho administration of chlorobutanol stimulates the synthesis of ascorbicacid, rouses the plasma level and restores fertility in the deficient animals

The reduction of the blood level of ascorbie acid and the occurrence of a scurvylike syndrome in rats on a vitamin A- and C-free diet suggest a functional relationship between these vitamins 217 Addition of ascorbie acid to the diet prevented or delayed the appearance of symptoms Simi lar observations have been reported in foxes *18 That ascorbic acid participates in the metabolism of other vitamins as well is indicated by its capacity for increasing the urinary excretion of citrovorum factor in rats and buman subjects 219 its influence on the storage of folic acid in the liver of chicks 120 and its effect of relieving deficiency symptoms of rats on a ration lacking thiamine, riboflavin, or pantothenic acid 221

Synergism between ascorbie acid and vitamin A has been noted in the rat as well as between C and E in the guinea pig Ascorbic acid enhances the growth promoting effect of vitamin A in the rat in a manner similar to tocopherol The mechanism may be through stabilization of vitamin A in the gastrointestinal tract The administration of vitamin E increases the storage of dietary ascorbie acid in the organs of the guinea pig

Sulzberger and Oser Proc. Soc. Exptl. Biol Med 32 716 (1934) See also Bundesen et al J Am Med Assoc. 217, 1692 (1941) McChesney et al J Pharmacol. Expli Therap 80 81 (1944)

kerap as 81 (1947) proc Soc. Erpt Bud Med. 47 374 (1948)
19 Frend and Willy Proc Soc. Erpt Bud Med. 47 374 (1948)
11 Langenecker Musellan Tully and King J. Bud Chem 127 4e3 (1939)
11 Langenecker Musellan Tully and King J. Bud Chem 127 4e3 (1939)
12 Langenecker Musellan Tully and King J. Bud Chem 16 313 (1948)
13 Welch tel J. Vatrison 35 (292 (1948))
14 Welch tel J. Pharmacol Erpt Throng 101 37 (1951) 183 403 (1951)
15 Welch tel J. Proc Soc. Erpt Bud Weld 75 130 (1954)

¹¹¹ Daft Fed Proc 10 350 (1901) McDaniel and Daft Fed Proc 10 387 (1901)

The ease with which ascorbic acid undergoes reversible oxidationreduction has suggested that the vitamin plays a role in cellular respiration. The oxidation of ascorbie acid is catalyzed by ascorbic oxidase. found in plant juices. The reverse reaction, the reduction of dehydroascorbic acid to reduced ascorbic acid, is catalyzed by ascorbic reductase (an enzyme found in plant juices) 222 in the presence of reduced glutathione. Though no similar enzyme systems have been observed in animal tissues, the administration of dehydroaseorbie acid to man is followed by increased urinary excretion of reduced vitamin C in an amount comparable to that which is excreted when the reduced form of the vitamin is fed. It is believed that this reduction occurs in the liver.

Certain in vitro findings lend ercdence to the view that ascorbic acid functions in vivo in oxidation-reduction reactions. Vitamin C plus minute amounts of iron cause a considerable increase in the oxygen uptake by phospholipides and by brain and liver suspensions. There is evidence. both in vitro and in vivo, that ascorbic acid functions as a coenzyme in the metabolism of tyrosine, although the exact nature of this action is not fully established. 323 A similar role is attributed to folic acid, the requirement for which appears to be increased in ascorbic acid deficient animals.

The action of urease, an enzyme which entalyzes the decomposition of urea to ammonia and carhon dioxide, is inhibited by vitamin C. This inhibition has been found to be due to the presence of small quantities of debydroascorbic acid, and is prevented by eysteine, which reduces the debydro form. Quinone, another oxidizing agent, can inhibit urease activity.

The recommended dietary allowances of ascorbic acid are shown on p. 1108. Approximately 1 mg. per day per kg hody weight is required for the maintenance of tissue saturation. These allowances are quite liberal, since clinical signs of scurvy do not appear even at considerably lower levels of intake. Considerable controversy exists as to the level of dietary intake of ascorbic acid necessary to insure adequate operation of the manifold functions in which this vitamin plays a role. Hence the recommendations of the National Research Council (p. 1108) provide for a substantial margin of safety. When healthy subjects have been subsisting on dicts containing the recommended levels of intake for prolonged periods, the administration of a test dose of ascorbic acid (400 to 1000 mg.) is followed in the next 24 hours by an increase in the urinary excretion of from 25 to 50 per cent of the test dose. At lower levels of intake for prolonged periods there is a tendency for the body to conserve the test dosc of the vitamin and lower excretion values are noted, indicating that body tissues are unsaturated.

No toxic symptoms are observed in man following the administration of large doses of ascorbic acid. One to 6 g. have been given orally and intravenously. Though ascorbic acid has been found in sweat, the amount lost through this channel even in excessive physical labor and in hot

¹²² Crook and Morgan Biochem J., 38, 10 (1944); Bukin, Biochimia, 8, 60 (1943). 222 Bessey , Lowe, and Salomon: Ann. Rev. Biochem , 22, 545 (1953).

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113 Bessey, Lowe, and Salomon: Ann Rev. Biochem, 22, 545 (1953)

climates is not significant when the dietary intake approximates the

recommended allowances Storage and Synthesis of Ascorbic Acid. As previously stated, all animal species other than man, monkeys, and guinea pigs, as well as lugher plants and many microorganisms are capable of synthesizing vitamin C The promptness with which scurvy develops when the susceptible species are deprived of vitamin C would seem to indicate the absence of a large store in their bodies. In man, a diminished intake of ascorbic acid results in a prompt fall in the blood level of ascorbic acid, though tissue stores may not yet be depleted Guinea pigs show histological symptoms of ascorbic acid deficiency after one week on a vitamin-C-free diet Higher concentrations of the vitamin are found in tissues of high metabolic activity-eg, the adrenal and pituitary glands and the intestinal wall. It is likely that the vitamin is present in these depots in order to meet tissue requirements, rather than for the purpose of storage Certain animals which synthesize vitamin C store it in their livers in sufficient amounts to meet their immediate requirements. The fruits and vegetables consumed by man as sources of ascorbic acid represent storage depots for plants which synthesize the vitamin

Distribution of Ascorbic Acid. 224 The outstanding sources of vitamin C are fresh fruits and vegetables Special mention may be made of cabhage, cauliflower, koblrabi, spinach, parsley, kale, broecoli, eresses, peppers, oranges, lemons, grapefruit, tangerines, limes, strawberries and gooseberries Other excellent dietary sources are cantaloupe, currants, papaya, persimmons, pineapple, asparagus, lima beans, green snap beans, and Brussels sprouts Unfortunately, much of the vitamin C originally present in foods is lost through processing so that for practical purposes the fruits and vegetables which are consumed raw contribute a major portion of the dietary ascorbic acid. For economic reasons, these are outside the reach of low income groups which obtain their vitamin C from the relatively inexpensive potato. During the germination of legumes as well as of cereal grains, marked synthesis of vitamin C takes place from an unidentified precursor within the seed Meat is low in vitamin C, although blood and glandular organs contain more than muscle Milk varies in its content according to the cow's ration and bence the season of the year Fresh raw milk is a good source of vitamin C, but most of it is lost in pasteurization evaporation, or dehydration Where milk is a major constituent of the diet, as it is for children, it is best to provide antiscorbutic protection from other sources, e.g., orange juice or syn, thetic ascorbic acid. The megaloblastic anemia observed in infants fed proprietary milk preparations has been explained as being due to a deficiency of folic acid secondary to chronic vitamin C deficiency The leaves and flowers of the gladiolus nettle hip, and paprika are exceptionally rich in vitamin C During World War II, rose hips green walnuts and wild cherries were used in Europe for the preparation of vitamin C

Chemistry of Ascorbic Acid. Ascorbic acid crystallizes in white

³¹⁴ For ascorbic acid values of foods see Appendix III

needles or plates having a melting point of 190° to 192° C. One g. dissolves in 3 ml, water, in 50 ml, absolute ethyl alcohol, and in 100 ml. glycerol. The vitamin is insoluble in henzene, ethyl ether, petroleum ether, and most organie solvents. Aseorhic acid possesses two asymmetric carbon atoms The vitamin has a specific rotation $|\alpha|_{0}^{20} = +21^{\circ}$ in water and +48° in methanol, p-Ascorbic acid, which differs from the L-isomer only in its configuration about the fourth carbon atom, is physiologically inactive. The acidic properties of ascorbic acid are due not to the carboxyl group, which is tied up in lactone form, but to ionization of the one group on the third carbon atom. The vitamin is a comparatively strong acid, as indicated by the acidie dissociation constants, $pK_1 = 4.17$ and pK2 = 11.57. A one-half per cent solution of ascorbic acid in water has a pH of approximately 3. At neutrality, the vitamin has an absorption maximum in the ultraviolet region of the spectrum at 265 mu which shifts toward the shorter wavelengths with decreasing pH. Ascorbic acid is precipitated by lead ion at nH 7.6, but the salt can be redissolved in mineral acid at pH 2.

Ascorbic acid crystals are stable in air for years In solution the vitamin is easily oxidized, the instability increasing with increasing pH. Mild oxidation such as that produced by air, hydrogen peroxide, ferrice chloride, quinone, iodine in acid or neutral solutions, or 2,6-dichlorophenol indophenol eonverts the vitamin to dehydroascorbic acid as shown in the reaction below. Dehydroascorbie acid retains the hiological activity of the vitamin and is reduced in animal tissues, a reaction in which sulf-hydryl compounds like glutathione play an important part. The reduction of the dehydroascorbic acid may be accomplished in vitro by means of hydrogen sulfade. Above pH 5 dehydroascorbic acid readily undergoes a

rearrangement in which the lactone ring is split. The product, diketogulonic acid, is no longer biologically active and is not reducible by
hydrogen sulfide. Theatment with hydrogen iodide, however, converts
this compound to dehydroascorbic acid, which may subsequently be
reduced with hydrogen sulfide to ascorbic acid. Once the lactone ring
has opened, the molecule readily undergoes further oxidation and may be
degraded to oxalic acid. The oxidation of ascorbic acid by molecular
oxygen is catalyzed by cupric and silver ions. Plant tissues contain
several enzymes, meluding ascorbic acid oxidase, polyphenol oxidase,

DETERMINATION OF ASCORBIC ACID

Chamical methods for the determination of vitamin C are based for the most part, upon the reducing properties of the vitamin. These procedures include titration of an acid extract with jodine, methylene blue in the presence of light, 2.6-dichlorophenol indophenol, or other oxidizing agents. Various acids have been used for the extraction, including acetic, trichloroacetic, metaphosphoric, and oxalic. The latter two acids serve not only to reduce the pH of the extracting medium, thereby stabilizing vitamin C, but also to form complexes with metallic ions. c g., copper, thereby preventing the catalytic oxidation of the vitamin. Titration with an oxidizing agent is not a specific reaction; other reducing materials interfere, e.g., stannous and ferrous sulfate, sulfhydryl comnounds, sulfides, thiosulfates, and reducing materials found in caramelized and fermented foods. Errors arise particularly in food products because of the presence of reductinic acid and reductones. Actually, reductone is hydroxypyruvic aldehyde, a compound formed by the alkaline hydrolysis of sugars. Reductinic acid is formed when pentoses are treated with acids. The terms reductone and reductinic acid, however, have been used rather loosely to signify other reducing compounds formed during heat-processing and storage of foods.

Numerous attempts have been made to increase the specificity of the oxidation-reduction methods, particularly that employing the 2,6-di-chlorophenol indophenol dye. A number of these are based on the fact that ascorbic acid reacts very rapidly with the oxidizing agent whereas many of the interfering substances react more slowly. The simplest expedient for increasing the specificity is to conduct the titration very rapidly, though it presents little improvement over the original method. A more satisfactory procedure involves measurement of the rate of reaction hy means of a photoelectric colorimeter. When observations are made five and ten seconds after mixing the ascorbic acid solution with the oxidant, and the values are extrapolated to "zero-time," the determination possesses sufficient specificity for vitamp C in most food products.

Modifications of the indopheaol titration have been aimed particularly at eliminating the interference of reductones and reductinic acid. These compounds are similar to vitamin C in structure, stability, and chemical reaction. Like ascorbic acid, their oxidative destruction is catalyzed by copper and by ascorbic acid oxidase. A number of procedures for climinating their interference involve condeasation of the reducing substances with formaldehyde, which eliminates the reducing properties of ascorbic acid, reductones, and reductinic acid; but the interfering compounds may be differentiated from the vitamin hy conducting the condensations at different pH levels. *** There is no doubt that these procedures increase the specificity of indophenol titration, though the interference of reductine acid is not completely eliminated. Morcover, errors arise because the procedures fail to take into account the presence of

³³¹ Lugg. Nature. 150, 577 (1942); Australian J. Exptl. Biol Med Sci., 20, 273 (1942); Mapson: Nature, 152, 13 (1943)

materials which affect the rate of combination of truscorbic acid with formaldehyde * 6

One method of improving the specificity of the determination of vitamin C involves the destruction of a corl ic acid with ascorbic acid oxidase thereby providing a blank correction value. This procedure however is laced on the assumption that the action of ascorlic acid ovida e is specific for vitamin C, whereas reductones are al o de-troyed 1 1 Natural pigments and stubborn turbs hites frequently interfere with the determination of vitamin C Several modifications of the indophenol methods may climinate these interferences since they involve extraction of the excess of unreacted dve with organic solvents (e.g. xvlene) im miscible with water 2 ?

Dehydroascorbic acid the rever- bly oxidized biologically active form of vitamin C does not possess reducing properties Since this form of the vitamin is frequently present in considerable proportion in proce-sed foods 210 it must first be reduced before it can be determined with oxidiz ing agents Hydrogen sulfide has been used most frequently for this pur nowe but this reagent has been enticized on the ground that it creates other reducing materials which interfere with the procedure. This objection is valid when simple titrimetric methods are employed for the determination of vitamin C However little interference arises from this source when the photometric method involving measurements at five and ten seconds is employed. Other methods for reducing the dehi droascorbic acid have been suggested but are not widely employed. These include the reduction of the oudized vitamin by a resting suspension of E coli²²¹ and by electrolytic reduction ²²

Dehydroascorbic acid and its mactive oxidation product diketogulonic acid couple with 2 4-dinitrophenylhydrazine to yield an osazone which gives a red color with strong sulfuric acid. By means of appropriate oxida tion and reduction reactions Roe and his associates bare adapted the dinitrophenylhydrazine reaction to the measurement of reduced and deby droascorbic acids (both biologically active) as well as diketogulonic acid. This permits estimation of the extent to which the original vitamin C content of foods has undergone oudation Since fresh or well preserved foods usually contain less than a per cent diketogulonic acid 124 the dinitrophenylhydrazine method is usually used in its simpler form adapted to the measurement of only the biologically active forms of

²²⁴ Cnow and Zilva Buchem J 37 629 (1943)

Sport and Lavra Beckers J at 1027 (1983)
 Tauber and Klemer J Boot Chem 118 9 9 (1973)
 Show and Zilva Boothem J 32, 199° (1973)
 Show and Zilva Boothem J 32, 199° (1973)
 Wat older and Okrent Z Physiol Chem 24, 22 (1974)
 Bukatech Z Physiol Chem 242, 29 (1979)
 See also Pepkowntz J Bod Chem 151

^{40. (1943)} The possibility of error due to the react on of anthory anna with the die is emphasized by Somers, & al. Science 118 17 (1992)

**Hothberg, Melnick, and Oper Ind E g Chm. Anal Ed. 15, 16° (1943) McMillan

^{**}Hocherty Steiner, san Off I had L. y. Larm and Ld. 15, 10 (1974) and Tolkunter Scarce 103 159 (1984)

1 Gunzalus and Hand J Bod Chem 141 8.3 (1941)

1 Gunther Booker, Z. 344, Z. (1932)

1 Roe and Kuether J Bod Chem 147 3.9 (1943) Roe and Octoberling Rod. 152.

J1 (1944) Roe et al Had., 174, 201 (1945

³¹⁴ Mills Damron and Roe Anal Chem 21 0" (1949

ascorbic acid. If oxidation during processing or storage is suspected, failure to account for discreptionic acid may lead to serious errors 255 356

The determination of ascorbie acid in blood and urine is employed in the diagnosis of clinical and subclinical vitamin C deficiency. These values fluctuate widely even in normal individuals depending upon the ascorbic acid intake just prior to the test. For this reason blood tests are conducted preferably when the subject is in a fasting state. A scrum or plasma level greater than 0.7 mg, per 100 ml, is regarded as normal whereas values less than 0.4 mg, are observed in source. Some clinicians set 1.2 mg ner 100 ml, as the minimum normal level. Individuals in a pormal state of nutrition excrete from 20 to 50 mg of ascorbic acid when fed a diet containing 100 mg, per day. Since both blood levels and urinary excretion values depend upon the ascerbic seid intake for one or two days prior to the test, a more satisfactory index of ascorbic acid nutrition may be obtained by measurement of the response of the subject to a large dose (500 to 1000 mg.) of ascorbic acid. Saturated subjects excrete anproximately 25 to 50 per cent of the test dose in 24 hours, and show a continued high blood level for hours following dosage. In one saturation test, an exerction of 20 mg, of ascorbic acid in the four-hour period following the ingestion of 200 mg is regarded as normal.

Plasma ascorbic acid values fail very rapidly when a deficient diet is consumed, though tissue stores including those of blood cells have not yet become depleted. It has been suggested that the ascorbic acid content of the leukocytes or of whole blood¹²⁷ is a better index of the state of vitamin C nutrition of the tissues than the blood plasma level. Oxybemoglobin interferes with the determination of ascorbic acid in whole blood by oxidation-reduction procedures unless it is saturated with carbon monoxide¹²⁷ or reduced ¹³⁸ The 2,4-dinitrophenylhydrazine method (see p. 1237) is suitable for determination of ascorbic acid in whole blood

One procedure intended for clinical diagnosis of ascorbic acid deficiency is based on intradermal injection of 2,6-dichlorophenol indopbenol followed by measurement of the rate of decolorization of the dvc ²¹⁹

Modified Titrimetric Method of Bessey: Principle. The method depends on the stochometric reduction of the dye 2,6-dichlorophenol indoplenol to a colorless compound by ascorbic acid. The titration is conducted in the presence of acetic and metaphosphoric acids in order to inhibit aerobic exidation catalyzed by certain metallic ions, to mactivate enzymes, and to precipitate proteins and hiberate protein-bound ascorbic acid. A modification of this method is based on the addition of an excess of the dye and measurement of the unreacted portion. In the presence of interiening pigmentation or turbulity, the dye can be removed by sylene extraction and rend in a photometer. It is necessary however, to guard against the extraction of sylene-soluble pigments present in some heat-processed foods, and certain non-vitamin-Csubstances (e.g., betanin of beets) which react with the dye.

³¹⁵ Penney and Zilva Biochem J . 37, 39 (1943)

Thoan and Gerjovich Science, 183, 202 (1946)
 Butler and Cushman J. Clin Investigation, 19, 459 (1940); Toe and Kuether. loc cit
 Kuether and Roe Proc Soc Ergl Biol Ved, 47, 487 (1941)

²³⁹ Reddy and Sastry Indian Med Gaz, 76, 476 (1941). ²⁴⁰ Bessey J Azoc. Official Agr. Chem., 27, 537 (1944).

¹⁴¹ See footnote 329, p 1234

Procedure Macerate an aliquot of the sample containing 5 to 50 mg ascorbic acid with 150 ml metaphosphoric-acetic acid solution. In a Waring Blendor Dilute to 200 ml and filter Tstrate a 10- to 100 ml aliquot with standard indophenol solution.

Preparation of Standard Indophenol Solution Dissolve 42 mg sodium hi carbonate and 52 mg sodium 1,6-dichlorophenol indophenol in 50 ml water Ditute to 200 ml Filter and store in the refrigeration not more than three days Dissolve 100 mg crystalline ascorhic acid in 100 ml metaphosphoric-acetic acid solution Dulute a 10 ml aliquot with 22 ml metaphosphoric-acetic acid solution, and titrate with indopnenol solution until the plac color persists for 5 seconds Calcinate and express the strength of the indophenol solution as mg ascorbic acid equivalent per ml regent Repeat the standardization each day with a freshly prepared standard ascorbic acid solution

CALCULATION Calculate the ascorbic acid content of the sample using the formula,

$\mathbb{T} \times S \times D = \text{mg. ascorbic acid per g. of sample}$

where I is mL of due used to titrate the unknown S is the standardization value expressed in mg ascorbic acid per ml and D is the dilution factor

Interpretation. The simple titrimetric method is applicable to the determination of ascorbic acid only in the absence of other reducing substances and where only the reduced form of the vitamin is present. Thus it may be applied to fresh orange grapefruit, lemon lime, or tomato juice and to some pharmaceutical preparations. For heat processed materials and other foods it is necessary to eliminate interfering substances and to determine the reversibly ordized, hiologically active form of the vitamin, deby droascorbic acid. Ferrous ion reduces the dye in the presence of metapho-phone acid. Pharmaceutical preparations containing reduced iron may be titrated in 8 per cent acetic acid solution free from metaphosphore acid. On the other hand, ferric ion interfers with the end point in the absence of metaphosphore acid hence the metaphosphoric acetic acid mixture should be employed as the titration medium when testing pharmaceutical preparations containing oxidized iron.

filtrate in a test tube, add 1 drop 10 per cent thiourea solution, and 1 ml. 2,4-dinitropheny lhydrazine reagent. ¹³⁴ Place the tube in a water bath maintained at 37° C. for exactly 3 hours. Remove and place in an ice-water bath along with a blank tube containing 4 ml. of "Norit" filtrate and 1 drop thiourea solution. To each tube in the bath add dropwise 5 ml. 85 per cent suifuric acid "1 with stirring. Finally add 1 ml. 2,4-dinitrophenylhydrazine reagent to the blank. Remove the tubes from the bath and allow to stand 30 minutes. Read the colors in a photoelectric colorimeter with a filter transmitting maximally at 540 ms, aetting the instrument at 100 per cent transmittance with the blank tube.

CALCULATION. Prepare a calibration curve by testing 4-ml aliquots of appropriate standards containing 0 25-150 µg ascorbic acid per ml carried through the entire procedure. Plot photometric density against µg assorbic acid per ml. From the curve, estimate the ascorbic acid concentration of the "Norit" filtrate of the unknown and multiply this by the dilution factor to obtain the ascorbic acid content of the sample

Interpretation. The 2,4-dinitrophenylhydrazine procedure measures total ascorbic acid. This may be partitioned into the reduced and dehydro forms hy including a simultaneous test in which the "Norit" treatment is omitted The latter procedure measures only dehydroascorbic acid The reduced ascorbic acid content is obtained by difference Reductones, degradation products of sugars, and diketogulonic acid interfere with the determination of ascorbic acid by this procedure

Determination of Ascarbic Acid in Whole Blood, Plasma, or Urine: Method of Roe and Kuether. 44

Procedure. Add 5 ml. whole blood or plasma, dropwise with stirring, to 15 ml. 6 per cent trichloroacetic acid in a 50-ml. centrifuge tube. Stir to obtain a fine suspension. Allow to stand 5 minutes, then centrifuge. Add 0.75 g. acid-washed "Norit" (see above) to the clear supernatant solution, and shake vigorously. Filter. Test 4-ml. aliquots by the 2,4-dinitrophenyihydrazine procedure as directed above.

To 2 ml. urine, add 38 ml. 4 per cent trichloroacetic acid solution. (A greater dilution should be made if the sample is expected to contain more than 300 mg. ascorbic acid per liter.) Add 1.5 g. acid-washed "Norit" (see above), shake vigorously, and filter. Test 4-ml. aliquots by the 2,4-dinitro-phenylhydrazine procedure as directed above.

Photometric Method of Hochberg, Melnick, and Oser. *** Principle. Ascorbic and is determined in the presence of other reducing substances by photometric measurement of the rate of decolorization of the dye 2,6-dieblorophenol indophenol Dehydrosscorbic and is determined after reduction by bydrogen sulfide

Procedure. Conduct all manipulations under an atmosphere of nitrogen and make all extractions and dilutions with solutions previously denerated with a stream of nitrogen. Homogenize the sample with an equal weight of 6 per cent metaphosphoric acid solution. 12 (Fresh vecetables and materials

³⁴⁶ Dissolve 2 g 2 4-dimitrophenylhydrazine in 100 ml 9 N sulfuric acid and fifter ³⁴⁷ To 100 ml distilled water, add 900 ml concentrated sulfuric acid sp. gr. 1 84

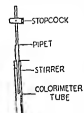
³⁶⁵ Roe and Kuether loc cut

¹⁴ Hochberg, Melnick and Oser Ind Eng Chem . Anal Ed. 15, 182 (1943)

²⁰ Metaphosphoric acid changes slowly in solution to orthophosphoric acid. The extractant should be stored in the refrigerator where it is stable for several weeks.

containing active oxidases should first be blanched by adding the sample directly to boiling 6 per cent metaphosphoric acid. After boiling for 5 minutes, cool and homogenize.) Dilute to a convenient volume with 3 per cent metaphosphoric acid. Shake mechanically for 15 minutes, then centrifuge. To 50 ml. of the clear supernatant fluid, add 14 ml. of citrate solution."

Measure reduced ascorbic acid in this allouor as directed below.



114 281 PHET USED FOR ADDITION OF AS-CORDIC ACID LATRACT TO DYE SOLUTION.

Convert dehydroascorbic acid to the reduced form by treating the metaphosphoric-citric acid extract with a slow stream of hydrogen sulfide for 20 mlnutes. Allow to stand for 2 hours, then flush out the hydrogen sulfide with a vigorous stream of wet nirogen (hubbled through water) for 2 hours.

Colorimetric Measurement, Determine the volume of extract necessary to decolorize 15 ml. 2,6-dichlorophenol Indophenol solution,¹⁸² To obtain optimal concentration for measurement in a photoelectric colorimeter, dilute this volume of extract to 30 ml. with metaphosphoric-cities cold solution ¹⁸²

Place a colorimeter tube containing 5 ml, of standard dye solution in the instrument, Add 5 ml, of the diluted extract from a rapid delivery piper***(Fig. 284) and measure the residual photometric density*** 5 and 10 secondo later. Determine the blank absorption of extraneous pigments in the extract by adding a few crystals of ascorbic acid to completely de-

Interpretation. This photometric method measures both reduced and dehydroascorbic acid. Appreciable concentrations of the latter are frequently found in processed and stored foods. The method is applicable to the determination of ascorbic acid in the presence of other reducing materials, which interfere minimally in this procedure.

RIOLOGICAL METHODS

Because of its nondependence on a dietary source of ascorbic acid. the rat cannot be used for vitamin C assay. The animal employed is the guinea nig, since it is extremely sensitive to the lack of this factor in the diet. The techniques used are based either on the determination of the minimum dose necessary for protection against or cure of gross scorbutic symptoms 356 or on the histological examination of the incisor teeth 357 after a test period of 14 days. The latter method requires equipment not always available in nutrition laboratories, though it has proved setisfactory for the determination of minimum protective dose where large enough groups are used to compensate for variations encountered. About twice as much ascorbic acid is required for protection of the dental pulp of guinea pigs as for prevention of gross scurvy. 358 The histological method has the advantages of high specificity and a short test period. Nevertheless, because of the special facilities required it has not been as widely used as the procedures based on prophylaxis or cure of gross symptoms.

The minimum daily allowance of a food which provides a guinea pie with complete protection from scurvy is a measure of its vitamin C potency. It has been suggested by Sherman, La Mer, and Campbell that by rating the nathological effects observed in the living animal and at autopsy, the fraction of complete protection afforded by subminimal doses may be estimated. While this method allows more animals to be considered in interpreting a given experiment, considerable experience is required in diagnosing and scoring the extent of the lesions.

ASSAY FOR VITAMIN C

Biological Method of Sherman, La Mer and Campbell. Guinea pigs, 6 to 8 weeks old and weighing from 250 to 300 g., are fed ad lib. the following scurvyproducing diet:359

⁵¹⁴ Sherman, La Mer, and Campbell J. Am Chem Soc. 44, 165 (1922); Coward and Kassner Brochem J , 30, 1719 (1936)

³⁷ Höjer Brit J Expli Path, 7, 356 (1926), Goettsch Quart J Pharm, 1, 168 (1928), Key and Elphick Biochem J, 25, 888 (1931), Fish and Harris Proc Roy. Soc London, 223, 489 (1933).

²³⁸ Eddy Am J Pub Health, 19, 1309 (1929).

¹⁵ Diets containing fat must be prepared fresh weekly, as guinea pigs reject rancid food Other scorbutigenic diets which have proved useful are

⁽a) Cambridge Nutration Laboratory (Harris et al.) Bran 80 oats 720 dried egg solk 40 salts 8 4, codliver oil 1

⁽b) Demole Z Vitaminforsch. 3, 89 (1934) Rulled cats 2 kg. dried whole milk 1 kg. (previously dried at 120° C for 2 hr). Mix and make into paste with whites of 6 eggs and water Mold into cakes 5 cm diam and bake 20-25 min on a tin greased with olive oil Supplement with 0 2 ml codhver oil weekly.

⁽c) Purma Rabbut Chow (Purma Mills, St. Louis, Mo.) produces scurvy in guinea pigunless supplemented with a source of ascorbic acid

Ground whole oats360	59
Heated nonfat milk solids set	30
Fresh butterfat162	10
Salt	1

This diet may he reinforced with yeast and codilver oil



Fig 285 Scorbette Guinea Pio Showing Scurry Position

In order to provide assurance of the health of the guines pigs and their capacity to grow, a short preliminary period may be included during which some source of vitamin C is fed, such as a leafy vegetable or orange juice



Following this period the food to be assayed is fed in graded doses as the sole source of vitamin C Feed 05 ml 01 per cent ascorbic acid or 5 ml orange or lemon juice to positive control animals as a supplement

In the absence of antiscorbutic vitamin, growth ceases in about 2 weeks, and a rapid decline in tic nextmortem findings are and to be more pronounced when the deficiency is not quite complete, because the lesions have time to reach an erapperated state. These findings include hemograpages (which may be subcutaneous, intramuscular, or intraabdominal), loosening of the teeth, fracility of the bones and teeth, and enlargement of the costochondral junctions of the ribs (beading)

Supplying a source of vitamin C to a guinea pig in even an advanced state of scurve usually will bring about prompt recovery, unless intercurrent disease has set in About 0.5-0.6 mg is required daily for complete protection OT CUITE

The vitamin C content is calculated from the minimum daily allowance which affords complete protection from scury, to a standard guinea nic. as above described. Comparison of this dose with the corresponding allowance of the crystalline L-ascorbic acid enables one to express the result of the bioassay on a me basis instead of in now obsolete International Units. The International Standard is crystalline L ascorbic acid, of which 0.05 me equals one International Unit of vitamin C

BIOFLAVONOIDS

In 1936, Szent-Gyorgy 364 prepared an extract from paprika and subsequently from citrus juice, which he claimed to have value heyond that of ascorbic acid in reducing capillary bleeding in man and in guinea pigs The name vitamin P was applied to this factor in the helief that it was a dietary factor necessary for the maintenance of normal capillary permeability Interest in various hemorrhagic conditions which do not respond to vitamin K or ascorbic acid has led to widespread investigation of concentrates high in the active permeability factor

Physiological and Clinical Aspects of Bioflavonolds, Although the literature is confusing on this point, a distinction should be drawn between capillary fragility caused by a change in the structural integrity of the capillary walls (e.g., loss of intercellular cement substance as in scurvy) and increased permeability without apparent tissue damage Clear differentiation is not possible in the usual tests based on subcu taneous petechial hemorrhage induced by negative pressure

Cumulated evidence has demonstrated the existence of a considerable

number of flavonoid or related compounds having the protective effect on capillary resistance associated with vitamin P activity. Some of the preparations used in animal and clinical studies, e.g., citrin, are actually mixtures of only partially identified compounds Effective doses vary over a wide range. In view of such considerations it is helieved that the activity of the flavonoids is a pharmacological one rather than a specific physiological function for which a dietary factor is essential Since continued application of the term "vitamin P" to one or another of these polyphenolic substances would lead only to confusion, it was recommended365 that the term be no longer employed The term "bioflavonoids" has been proposed for this class of compounds 366

Scarborough 167 describes the characteristic syndrome of petechial 344 Szent Gyorgy and Rusznyák Nature 138 "7 (1936)

³⁴⁴ Joint Committee on Nomenclature American Society of Biological Chemists and American Institute of Nutrition Science 112 628 (1950)

^{***}Oser cited by Bryant J 4m Parm Assoc 39 480 (1950)

**Scarborough et al. Buchem J 33 1400 (1939) Lancet 2 610 (1938) 644 (1910)

£dinbr ph Ved J 50 85 (1943)

bleeding as accompanied by pain across the shoulders and in the legs, lassitude and undue fatigue, low capillary resistance, slightly prolonged bleeding time, and low serum calcium. The biofilar onoids are effective in relieving this syndrome but nurs ascorbic acid is not

It has been postulated by the French workers¹⁶ that epinephrine is the compound directly responsible for the maintenance of capillary integrity and that the protective action of the bioflav onoids is an indirect one, based on their inhibitory effect on the oxidation of epinephrine (demonstrable both in ziro and in ziro). Certain flav onoids have been shown to produce a fall in afternal blood pressure in animals³⁴⁹ and to inhibit hyaluronidase¹⁷⁰ and histidine decarboxylace,²⁷¹ but whether these effects are causally related to capillary resistance, and how, have got been established.

There is conflicting evidence in the literature regarding the clinical value of these compounds, which probably results from the fact that there are many factors responsible for the maintenance of normal capillary resistance. Clinical methods of measuring capillary fragility are inadequate because of diurnal variations and differences in subcutaneous capillary helds in different areas of the body. The dosages employed clinically depend upon the purity of the extracted flavones, the crystalline glycosides have been administered in divided dosage in the order of 50 to 150 mg per day.

Occurrence of the Bioflavonoids. Citrin or similar gly cosides baving hioflavonoid activity have been isolated from paprika, juice or peel of citrus fruits, buckwheat, black currants, cured tobacco, and the leaves of a wide variety of plants.

Chemistry of the Bioflavonoids Most if not all of the active flavonoids occur in nature as glycosides, either alone or in combination Citrin from lemon peel is said to be a mixture of the flavonone glycosides, hesperdin and enodictin (the corresponding aglycones being hesperetin and enodicty of, respectively) although its exact composition has not been clearly established. The relation of flavanones and flavonols to the parent nucleus of flavone (2 phenyl-1,4-benzopyrone) is shown by the following structural formulae.

Flavonoids	Nucleus	Substituents	Positions
Eriodiety ol*	Havanone	4 (011)	5, 7, 3', 4'
Hesperetin*	Flavanone	3 (OH)	5, 7, 3'
•		1 (OCH ₂)	1'
Hesperidin	Flavanone	2 (OH)	5, 3'
•	1	1 (OCH ₂)	4'
		Glucorhamnose	7
Rutin	Flavonol	4 (OH)	5, 7, 3', 4'
		Glucorhamnose	3
Quercetin*	Flavonol	4 (OH)	5, 7, 3', 4'
Quercetrin	Flavonol	4 (OH)	5, 7, 3', 4'
-		Rhamnose	3

* Agly cones

Certain chalcones, i.e., compounds resulting from the opening of the pyrone ring at the 1-2 position, are active, as are also catechin (not to be confused with the dihydric phenol catechol) and its stereoisomer epicatechin.

Opened ring of chalcone nucleus (cf. Flavone)

Catechin

Determination of Bioflavonoids. The varied chemical structure of naturally occurring bioflavonoids and their active derivatives has militated against the possibility of a chemical assay correlated with the specific type of biological activity However analytical methods have been described for flavonoids in certain plant or animal tissues. The biological assay is based upon the observation of petechial hemorrhages under controlled conditions similar to the procedure employed for the diagnosis of "vitamin P" deficiency in man. Bacharach, Coates, and Middleton? have described a guinea pig assay in which crystalline hesperidin is used as the reference standard. The critical petechial pressure is determined from the negative pressure required to produce hemorrhages when a suction cup, furnished with a glass window for visibility, is applied to the prepared area of skin.

VITAMIN D

Early in the history of vitamins, antirachitic activity was ascribed to fat-soluble vitamin A, but the work of McCollum and his collaborators²⁷³

⁴⁷² Bacharach, Coates, and Muddleton Biochem. J., 36, 407 (1942); See also Bourne: Nature, 152, 659 (1943).

²¹³ McCollum, et al.: J. Biol. Chem , 50, 5 (1922); 51, 41 (1922); 53, 293 (1922).

established the distinction between the growth-promoting, antiophthal mic vitamin A and the antirachitic factor to which the name vitamin D was assigned. The specific role of this vitamin in the prevention and cure of rickets (rachitis) is integrated with the metabolism of calcium and phosphorus, in fact, in some species, e g , the rat, the absence of vitamin D from the diet does not result in rickets unless the normal dietary ratio of Ca P is disturbed A condition analogous to human rickets may be produced experimentally in this species by feeding a diet high in calcium, low in phosphorus, and free from vitamin D On a low-calcium, highphosphorus diet, the bones develop an osteoporotic condition resembling osteomalacia The function of vitamin D appears to be to mobilize these calculving elements, so as to make possible their most efficient utilization even when the dietary supply is inadequate or disproportionate. It has been shown that even if calcium and phosphorus are fed in proper quantity and ratio, the absence of vitamin D, especially if prolonged, will result in poorer calcification of the bones than obtains under normal conditions 374

Mild cases of rickets are often unrecognized, though they are quite prevalent, and may not have serious consequences until later years. The work of Jeans and his associates has emphasized that the requirement of vitamin D for optimizing month exceeds the antirachitic requirement. Rickets in its more severe manifestations is seen more frequently among urban children of poor economic status. In regions where sunshine is prevalent and customs permit bodily exposure the disease is of rare occurrence. In India rickets is less common among the poor who live an outdoor existence than among the wealthy who observe the rite of

seclusion (purdah)

Physiological and Clinical Aspects of Vitamin D. Rachitic bones are characterized by a lower content of mineral matter and an apparent overgrowth of esteoid cartilage in consequence of which normal rigidity is absent The sequelae of this condition in children are misshapen bones (e.g., bowlegs), epiphyseal chargement (e.g., knock-knees and beading of the ribs, the "rachitic rosary"), delayed closure of the fontanelle, retarded cruption of the teeth, disturbances of respiration due to deformity of the thoracic cavity (pigeon breast) etc. In later life, difficult childbirth may be encountered because of pelvic deformities. The changes in bone composition are illustrated in the table on p. 1245.

The calcium phosphate compound in all bones, normal or rachitic, appears to be Ca₁(PO₂), which in the normal adult rat is accompanied by CaCO₃ to the extent of about 15 per cent of the total calcium *** The ratio of tertiary phosphate to carbonate is reduced in the rachitic animal From evidence based parily on solubility product studies it has been suggested that primary calcification differs in composition from old

Composition of Normal and Rachitic Bones (Schabad)376
(In fer cent of dry fat-free matter)

В	one	II aler	Ash	Organic Matter	Calcium	Phos phorus
7.1	Normal	14 4 32 9	40 2-46 6	26 9-39 1	15 5-18 1	5 4 8 3
Rıb	Rachitic	42 4-66 4	7 9-32 0	20 7-22 4	3 0-12 0	1 4 5 6
	Normal	13 0-16 1	47 6-51 7	32 2-36 5	18 8-19 9	7 9-9 0
Occiput	Rachitic	29 0-35 9	34 3-40 6	26 1-31 6	13 6-17 2	6 0-7 8

bone ²⁷⁷ Refractive index and x-ray studies show that the solid inorganic phase of bone consists of minute crystals which are normally oriented lengthwise along the bone, but in rickets the cement substance (collagen) is broken down and the x-ray diffraction pattern indicates a discriented crystalline structure ²⁷⁸

Many suggestive facts are known, but no complete explanation of the mechanism of vitamin D activity is available. The diminished retention and altered paths of elimination of calcium and phosphorus in rickets (illustrated in the table on p. 1246) provide the basis for the conception that vitamin D acts by promoting the absorption of calcium and phosphorus from the small intestine A causal relation between the intestinal pH (probably only in the cecum and colon) and the retention of the calcifying elements is suggested by the greater alkalimity (and bence the precipitation of calcium salts) in rickets, but it is not known which of these is the determining factor and, if it is the former, what ultimate mechamism is responsible for its alteration in rickets. While the effect seems to be mediated through an increase in permeability, vitamin D does not appear to influence the excretion of calcium into the intestine, except indirectly Vitamin D is claimed to increase gastric secretion and thus facilitate calcium absorption. The shalper peaked blood sugar tolerance curve after vitamin D administration is also suggestive of an effect on intestinal absorption

A significant agent in ossification is the enzyme alkaline phosphatase ³⁷⁸. This enzyme occurs in bone, ossifying cartilage, the hidneys and intestinal mucosa, and in low concentration (5 to 15 units per 100 ml) in blood. By hydrolyzing phosphoric acid esters in bone it liberates inor-

^{*} Frepared by Orgler (Ergebnisse inn. Med. Kinderheill. 8, 142 (1912)) from the data of Schabad (Arch. Kinderheill. 52, 47 (1909), 53, 380 (1910), 54, 83 (1911)) and cited by Hess (see Ibbliography p. 1296).

Hess (see Libbography p 1296)
"I kramer and Shear Proc Soc Exptl Biol Ued 25 141 and 285 (1927 28)
"I Taylor and Sheard J Biol Chem 81 4 9 (1929) Clark and Mrgudich Am J
Physiol 108 74 (1934)

Propose 108 4 (1934)

11 Robison et al. Biochem J. 17 286 (1973) 18 "40 and 1354 (1974) 19 153 (1975)

20 847 (1976) 23 767 (1929) 24 1922 and 1927 (1930) hay Biochem J. 20 791 (1976)

21 25 55 (1978) J. Biol. Chem. 89 235 and 249 (1930)

game phosphate which modifies the calcium and phosphate ion concentrations to the extent that the solubility product is exceeded and excess calcium phosphate is deposited. The optimum pH range of alkaline phosphatase is 8 4 to 9 ±

The fact that phosphatase is concentrated in the ossifying centers of bone cartilage even in rickets would tend to indicate that in this disease

TABLE SHOWING RETENTION AND ELIMINATION OF CA AND P IN RACHITIC RATS AND DURING HEALING*

(AVERACES OF FOUR RATE FXPRESSED ON THE BASIS OF ONE RAT PER PERIOD)

		Calcu	unt			Phosph	torus	
Lernods		Distribution (per cent of intal e)				Distribution (per cent of inial c)		
Intal e mg	Intal e mg		Excreted		Intal e	2	Eza	rete I
	Relained	Urine	Feces	Pelained		Urine	Feces	
I (R ekets)	365	15 6	12 1	72 3	105	36 2	0.0	63 8
II (Healing)	163	47 2	0.7	52 1	159	39 6	20 5	39 !

^{*}From unpull aled data of ta ned in the sen or author a laboratories. S milar observa t one are reported by St ohl. Bennett. and Weed. J. Biol. Chem. 79. 207 (1928) Per od. I. A. 6-day, period in the fourth week of a h. gh. Ca. low P rachitogen dict

Per od II A 5-day per od dur ny the succeeding week after the Ca P ratio was d min ist ed by autst tuting CalifO. 2lifo for CaCO:

the supply of substrate may be at fault. In nickets there is not only over production of osteoid matrix but also proliferation of cartilage cells which are an abundant source of phosphatase. The small amount of this enzyme normally found in plasma is greatly increased in generalized boson desorders, probably by diffusion from the osseous tresus (see p. 253), in which a compensatory stimulation of osteoblastic activity occurs it has been suggested that two mechanisms operate in bone formation, the phosphatase mechanisms which produces in the bone matrix fluid a condition of super-sturation with respect to bone phosphate and an 'inor game mechanism which favors deposition of this salt from super saturated solitions

New light has I cen east on the I tochemical role of alkaline phosphatase in rickets by the work of Letterstrom and I junggren ¹³⁰ who studied the action of the water soluble phosphorilated derivative of vitamin D (D₂P). They showed that D₂P has a remarkable capacity to activate phosphatase especially from bone. It would appear therefore that vitamin D may function as an activator of alkalite phosphatase not only in the indexina an option of the phosphatase is only in the formal to the phosphatase in the following the phosphatase is only in the following the phosphatase in the following the phosphatase is only in the following the phosphatase in the following the phosphatase is only in the following the phosphatase in the following the phosphatase is the following the phosphatase in the following the phosphatase is the following the phosphatase is the following the phosphatase in the following the phosphatase is the following the phosphatase is the phosphatase in the phosphatase is the phosphatase in the phosphatase is the phosphatase in the phosphatase is the phosphatase in the phosphatase is the phosphatase in the phosphatase is the phosphatase in the phosphatase is the phosphatase in the phosphatase is the phosphatase in the phosphatase is the phosphatase in the phosphatase is the phosphatase in the phosphatase is the phosphatase in the phosphatase is the phosphatase in the phosphatase in the phosphatase is the phosphatase in the phosphatase is the phosphatase in the phosphatase is the phosphatase in the phosphatase in the phosphatase is the phosphatase in the phosphatase in the phosphatase is the phosphatase in the phosphatase in the phosphatase is the phosphatase in the phosphatase in the phosphatase in the phosphatase is the phosphatase in the phosphatase in the phosphatase in the phosphatase in the phosphatase in the phosphatase in the phosphatase in the phosphatase in the phosphatase in the phosphatase in the phosphatase in the phosphatase in the phosphatase in the phosphatase in the phosphatase i

tubular reabsorption, all of which are involved in clinical rickets. A more general role for vitamin D in earbohydrate oxidation is suggested by in ritro studics381 in which D2P was shown to have an activating effect on the overen untake of kidney mitochondria. The respiratory enzyme complex here involved is known to be concerned in the tricarbovylic acid cycle and in fatty acid oxidation.

Efforts have been made to correlate the action of vitamin D with the important role played by the parathyroid hormone in the maintenance of calcium equilibrium in the serum. Hyperparathyroidism is accompanied by high serum calcium, and apparent dissolution of this element from the bones which, when prolonged, may lead to osteomalacia. Parathyroid extract does not cure rickets. Though there are similarities in the effects of vitamin D and parathyroid hormone on calcium mobilization, evidences of differences in their mode of action-e.g., in tetany or when given in overdosage-make doubtful a direct hormonal role in vitamin D activity.

The vitamin D requirement of man is influenced by age, color, pregnancy, lactation, the mineral content of the diet, and conditions which affect intestinal absorption. The vitamin D needs of adults and older children may be satisfied by sufficient exposure to ultraviolet radiation. According to the National Research Council, the recommended daily allowance of this vitamin is 400 units daily, except during pregnancy. lactation, and the first year of childhood, when it is increased to 800 units See p. 1108 Vitamin D may be administered not only subcutaneously or intraperitoneally 382 but by direct absorption through the skin. 282 although the dosage requirements under these conditions are not well defined.

Even when massive doses are given, the transfer of vitamin D to human or cow's milk is limited to only a few per cent The administration of 40,000 units per day during early lactation of mothers increased the vitamin D content from very low or zero levels to a range of 125 to 583 units per liter. Vitamin D milk of commerce is produced by direct addition of irradiated sterols or concentrates; formerly "metabolized vitami D milk" was obtained by controlled dosage of cows.

Some success has been claimed for the more or less empirical use o large doses of vitamin D in the treatment of arthritis, but the reports it the literature are conflicting and the physiological mechanism involved i not known

The belief that excessive doses of vitamin D were toxic owed its origin in part to unfortunate experiences with an early German product "Vi gantol," which probably contained an excessive proportion of toxistero (see p. 1252) due to overirradiation of the sterol. Later studies have shown that the margin between the therapeutic and the minimum toxic dose of vitamin D is very wide. Several hundred times ordinary therapeutic doses

Zetterstrom: Acta Chem. Scand., 5, 343 (1951)
 Soames Biochem J., 18, 1349 (1924); Hess, Weinstock, and Helman J. Biol. Chem. 63, 305 (1925); Kraimer, et al. J. Biol. Chem., 71, 699 (1927)
 Hume, Lucas, and Smith: Biochem. J., 21, 362 (1927). Helmer and Jansen: Studies

Inst. Din Thorac, 1, 83 (1937).

must be administered daily for several weeks before toxic effects are noted. It has been suggested that infants be given doses of 250,000 units twice a year for protection against nickets. The chief symptoms of toxicity due to overdosage of vitamin D are anorexia and polyuna. Calcification of soft tissues, and particularly of the renal arterioles and the aorta, are observed in advanced stages, although hypertension is not encountered Vitamin D overdosage is indicated by a rise in serum calcium. The toxic effects disappear upon discontinuance of the dosage.

The calcium content of the blood serum of rachitic children is normal (0-11 mg per 100 ml) or only slightly lower, but the inorganic phosphorus content may be reduced to balf the normal values of 4-5 mg, or even lower in severe cases. There are seasonal variations in the blood phosphate values, the minimum occurring in late winter and early spring, and the maximum in midsimmer. These changes run parallel to the scasonal variation in solar ultraviolet radiation and coincide with the variations in the incidence of clinical nickets. They may also account for the spontaneous healing often observed in summer.

These observations as well as the well-known geographical distribution of rickets, have been attributed to the effect of exposure to sunlight. The therapeutic value of the sun's rays in rickets was noted by Huldschinsky, who also studied the use of artificial sources of ultraviolet radiation [Certain of these rays cause a synthesis of vitamin D from a precursor [7-dchydrocholesterol] in the skin, and a similar synthesis can be produced in animal and plant oils from this or other provitamins.

For further discussion of the clinical aspects of vitamin D, see the

American Medical Association syllabus, p. 1292

Ultraviolet Radiation. The visible region of the spectrum comprises only a small portion of the entire scale of radiant energy. The colors in the visible spectrum result from differences in the frequency—or inversely, the wavelength—of the radiations. In the invisible regions of electromagnetic vibrations, there are the longer radiations (infrared or heat rays, Hertzian, and radio waves) and the shorter radiations (ultraviolet rays, gamma rays, v rays, etc.) The relation between these forms of radiant energy is illustrated in Fig. 287

The antirachitic region of the spectrum extends from about 256 m_µ to about 313 m_µ, whereas the shortest solar radiation is about 290 m_µ. It is only when the atmosphere is free from smoke, fog, dust, clouds, etc that solar energy of wavelength shorter than 300 m_µ reaches the earth, and even then its intensity is very slight. On a clear day at sea level the energy distribution is approximately 1 to 2 per cent ultraviolet, 42 to 53 per cent visible and 57 to 63 per cent intrared. The himted antirachitic protection offered from this source helps explain the frequency of rickets and the need for dietary sources of preformed vitamin D. Ultraviolet reduction exerts its maximum activating power at the wavelength of maximum absorption by the provitamin, e.g., at 281 m_µ (range = 275 to 300 m_µ) in the case of crosterol. The energy required for activation is 75 × 10¹¹ quanta per unit of vitamin D.

Ultraviolet energy emanates from incandescent solids. The distribution and intensity of this energy depends on the nature of the incandescent

source. The most widely used sources of ultraviolet radiation in experimental or therapeutic work are the carbon are and the quartz mercury are. Carbon are lamps are provided with carbons containing metallic cores of definite composition. The quartz mercury-vapor lamp consists of a quartz chamber in which an are discharge takes place hetween electrodes of mercury or of mercury and tungsten. The ultraviolet component of the radiation from these lamps depends on the energy input and, in the case of the mercury vapor are, diminishes with the age of the lamp.

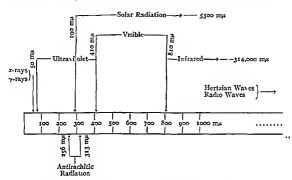


FIG 287 CHART SHOWING DISTRIBUTION OF RADIANT ENERGY.

The figures indicate the wavelengths, expressed in milhmicrons. 344

Ordinary window glass is opaque to ultraviolet radiation but quartz is transparent Various makes of glass of special composition permit the transmission of antirachitic radiation and are used in schools, hospitals, solaria, etc. Certain plastic films are also transparent to the antirachitically active rays, and these have been used in windows of poultry houses

The variation in distribution and intensity of ultraviolet radiation from the sun or from artificial sources makes necessary some means of measurement. No satisfactory method has been developed for analyzing the antirachitic region alone, and of the biological and photochemical methods it may he said that no two of them measure exactly the same region and none of them gives an index of the distribution of energy. The only accurate methods are those based on a physical measurement of the energy, such as those employing a thermopile and galvanometer. Radiation of any region of wavelength, selected by optical filters or a quartz monochromator, may he absorbed by a thermocouple, causing a flow of current which can be measured by a galvanometer previously standardized

³³⁴ One micron (μ) = 0 001 mm.; one milimicron ($m\mu$) = 0 001 μ ; one Angstrom unit (Λ) = 0 1 m μ

against radiation of known intensity. Such a method, while nonselective and reliable, is beyond the scope of most biological laboratories

Of the photochemical methods³⁵⁵ there may be mentioned (1) Clarl's, in which the number of minutes required for a paste of lithopone, or preferably pure zine sulfide, to darken to a definite shade is taken as the index and expressed as lithopone or ZnS units of ultraviolet energy, (2) Poble's in which the index is the time required for the ultraviolet energy to produce a blue color in a solution containing potassium iodide, sodium throsulfate, and starch (3) Webter, Hill, and Endinov's in which the fading of the blue color in an acctone—methylene blue solution is measured by comparison with known standards, and finally (4) the method of Anderson and Robinson, given below

Photochemical Method for Measuring Ultraviolet Intensity Principle In the presence of a uranium salt which acts as a catalyst a solution of exalic acid is decomposed by ultraviolet but not by visible radiation. The degree of decomposition is determined by titration with pota-sum permanganate.

Procedure A solution containing exactly 6.3 g. c.p. oralic acid (H.C.O. 211,0) and 4.27 g. uranyl sulfate (UO,SO.311,0) per liter of distilled water is prepared and kept protected from the light Place 25 ml of this solution in a fused quartz cell 3.35 × 10 cm in area and 1.65 cm thick **The unshielded area of the cell is exposed directly to the source of ultravolet energy for a definite period. In the case of solar measurements a one hour exposure is used the angle of the cell being changed every 5 minutes to keep the rays perpendicular to the surface.

After the exposure the contens of the cell are runsed into a flash with 50 ml distilled water, and 2 ml concentrated sulfuric acid are added. The solution is then boiled and titrated while hot with a solution of potassium permanga nate standardized to be equivalent to the oxalic acid solution (theoretically 3 16 g. NinO, per liter).

"CALCULATION" (25 - ml of f.MnO, used) × 0.63 ≈ mg oxalic acid decomposed Duplicate determinations sloul agree within 0.06 mg. The maximum value found by Tonnex for soul git at Cheago was 7.12 mg per hour Sunburn of untanned skin (the crythema reaction) was observed whenever the intensity exceeded half this maximum value.

Storage of Vitamin D. The animal hody is able to maintain a reserve of vitamin D, the amount depending on the dietary supply and on the event of exposure to the synthesizing influence of ultraviolet radiations. The rate of depletion of the bodily store of vitamin D, when a supply of this factor is lacking, is affected by the ratio of valenum-to phosphorus in the diet which under normal conditions should be between 1, 1 and 2.1

Young animals acquire some degree of autirachitic protection by virtue of the transfer of vitamin D into the mother's milk although the efficiency of this transfer is very low (about 1 to 2 per cent). By feeding fish liver oil or irradiating the mother during the periods of gestation and lactation the vitamin D notency of the milk can be increased somewhat, but ordinarily both cow's milk and human milk are poor sources of the antirachitic vitamin. The fortification of cow's milk with vitamin D is discussed on n 1256.

Chemistry of Vitamin D. Five years after Huldschinsky's 287 demonstration of the therapeutic value of ultraviolet irradiation on rickets. Hess³⁵⁸ and Steenbock²⁸⁹ in 1924 independently revealed the discovery that antirachitic activity could be induced in foods by exposure to ultraviolet radiation. It was soon demonstrated that the antirachitic factor was contained in the nonsaponifiable fraction of activated oil, just as in the case of codliver oil. Interest then centered on the activability of phytosterol and cholesterol/ the important unsaponifiable lipides of plant and animal tissue, respectively. At first it appeared that cholesterol was the specific precursor of vitamin D, but later work revealed other sterols to function in this role, viz., ergosterol in the lipide fraction of plants, yeasts, and fungi, and 7-dehydrocholesterol in animal sources. Ergosterol had originally been isolated from ergot, the fungus of rye, Following the discovery of its provitamin nature it was extracted commercially from yeast, 7-Dehydrocholesterol was originally obtained from brain tissue, crustaceans, mussels, and other shellfish, but is now produced synthetically. The activated product of ergosterol is called vitamin D: or calciferal, whereas that of 7-dehydrocholesterol is vitamin D.

From the structural formulas it will he seen that activation of these sterols involves opening of the B ring and conversion of the methyl group

between the A and B rings to methylene.

The synthesis of vitamin D from ergosterol is independent of the wavelength of the ultraviolet radiation over the range of selective absorption (about 256 to 313 mμ), but is a function only of the absorbed energy. 390 It is therefore possible to activate ergosterol by exposure to the unobstructed rays of the sun in spite of the fact that radiation of such short wavelength, even under ideal conditions, hardly ever exceeds 0.1 per cent of the total solar energy. 391 The efficiency of solar irradiation may be affected by a relative preponderance of inactive decomposition products produced by the longer waves 392 Activation is also induced by means of electronic streams, x-rays, radioactive emanations, and high-frequency alternating current.

³⁵⁷ Huldschinsky: Deut med. Wochschr , 45, 712 (1919). 388 Hess. Am. J. Diseases Child., 28, 256 (1924); Hess and Weinstock. J. Biol. Chem.

³⁵⁴ Steenbock and Black: J. Biol. Chem , 61, 405 (1924).

¹⁰⁰ Kon, Damels, and Steenbock: J. Am Chem. Soc., 50, 2573 (1928); Webster and Bourdillon: Biochem J., 22, 1223 (1928); Nature, 123, 244 (1929); Marshall and Knudson. J. Am Chem. Soc., 52, 2304 (1930).

Coblentz, Dorcas, and Hughes: Bur. Standards Sci Paper No. 539, 21, 535 (1926); J. Am Med, Assoc, 88, 390 (1927); Greeder and Downes, Trans Illum, Eng. Soc., 25, 378

^{(1930);} Forsy the and Christison: J. Optical Soc. Am., 20, 396 (1930).

111 Lahousse and Gonnard, cited by Bills Physiol Revs., 15, 1 (1935).

In the ultraviolet activation of ergosterol a series of products is formed depending on the wavelength of the radiation, the duration of exposure, overheating the nature of the solvent, the presence of oxygen, etc The changes induced in the absorption spectrum of ergosterol by irradiation

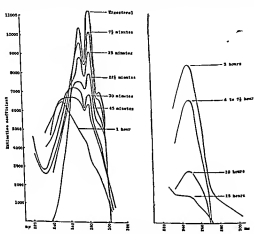


FIG. 288 1 PFICT OF PROLONGED IRRADIATION ON THE ULTRAVIOLET ABBORP TION SPICTREM OF LEGORIFROL. Courtesy Bid: Blookyedi and Got J Bid Chem 80 557 (1928)

are shown in Fig. 288. The following scheme summarizes the sequence in which the isomeric products of irradiation appear.

$$\begin{array}{c} \text{Lrg we rol} \rightarrow \text{Lumisterol} \rightarrow \text{Tachy sterol} \rightarrow \text{Calciferol} \\ (\text{Uniform a Di}) \rightarrow \text{Calciferol} \\ \end{array}$$

methylene A similar change occurs in the activation of 7-dehydro-cholesterol (p. 1254)

The properties of ergosterol, 7-dehydrocholesterol, and their respective activation products are shown in the following table

	Ergosterol	Vilamin D ₂ (Calciferol)	7 Dehydrochol esterol	Vitamin Di
Formula	C11H (1OH	C28H42OII	C27H42OH	C27H42OH
Double bonds	3	4	2	3
Melting point	166° C	116° C	143° C	83° C
Digitonin	Precipitate	No precipitate	Precipitate	No precipitate
Absorption bands	260 270 282 293 5 m µ	265 mµ	Same as ergosterol	Same as calciferol
Optical rotation	In CHCl ₃ $[\alpha]_D^{10} = -132^{\circ}$	In C H ₅ OH $[\alpha]_D^{to} = +102.5^{\circ}$	In CHCl ₁ $[\alpha]_D^{70} = -113.6^\circ$	In acetone $[\alpha]_D^{20} = +833^\circ$
Vitamin D potency rats chicks	None None	40 000 000 u /g 400 000 u /g	None None	45 000 000 u /g 45 000 000 u /g

The demonstration*** that activated ergosterol when fed to chicks on an equivalent rat-unit basis is only about one hundredth as effective for them as codiver oil paved the way for studies proving that cal ciferol is not identical with naturally occurring utamin D. By means of the comparative chick vs. rat assays, Waddell*** showed that the proving in impure cholesterol was not ergosterol, since the relative species effectiveness of irradiated cholesterol is similar to that of the instruction of the comparative which can acquire antirachite potency upon irradia.

Massengale and Nussmeier J Biol Chem 87 423 (1930)
 Waddell J Biol Chem 105 711 (1934)

tion, 72 including in addition to the natural precursors ergosterol and 7-dchydrocholesterol, 22-dihydrocysterol, 7-hydrocyscholesterol, and 7-dehydrocytosterol linere is also exidence for the chemical activation of cholesterol (by forming cholesterylene sulfonic acid) and of ergosterol (by treatment with intrite)

The products of irradiation of 7-deliydrocholesterol are analogous to those of ergosterol, the antirachitic substance being identical with the naturally occurring vitamin in fish liver oils 7-Deliydrocholesterol can be sufficied by mammals and is present in subcutaneous tissue Chemically it differs from ergosterol oils in the side chain at position 17, which in this case is saturated and has one less CH₂ group. The initial ultraviolet absorption curves and the transitional changes during irradiation of ergosterol and 7-dely drocholesterol are identical.

Distribution of Vitamin D. The most abundant natural source of vitamin D is codliver oil. Nevertheless, whereas ordinary codliver oil possesses a vitamin D activity of approximately 100 to 200 U.S.P. units ner gram, the liver oils of certain other species of fish may be many times more potent. 399 Some liver oils which are blended for medicinal use are high in vitamin D, eg, those from various species of tuna. Other oils, eg, those of the swordfish and halibut, are also high in vitamin A. (See table n 1115) The vitamin notency of fish liver oils is affected by snowning (and hence the season of the catch and the oil content of the livers) and by the age of the fish. Varying amounts of vitamin D are also found in the hody oils of many marine animals, among them salmon. herring, sardine, shrimp, oysters, etc. 400

Whether fish are able to synthesize vitamin D, as indicated by the experiments of Bills. 199 or whether it is derived from the caplin and other forms of food they consume, has not been definitely settled. There is evidence 101 that the vitamins of codliver oil are derived indirectly from unicellular marine organisms via copepods, larval decapods, and mollusca present in marine plankton, which in turn is consumed by the cod However, Leigh-Clare 102 failed to demonstrate the presence of vitamin D in these diatomaceous organisms, although other investigators report the presence of the vitamin in marine plankton 403 especially during the summer months when it is abundant near the surface of the water.

When the livers of cod are in the process of becoming spent, the vitamin D potency of the oil hears an inverse relation to the oil content 404 This has been attributed to the relatively rapid depletion of the glycerides during the starvation period, leaving behind the more resistant antirachitic factor. In a large-scale experiment conducted at the Norwegian fishing grounds, however, the senior author found that at the spawning season, when the livers were rich in oil, there was no demonstrable difference in antirachitic potency between the oil of male and female cod The hyers of the females had a somewhat lower oil content but were larger than those of the males 400

No increase in the vitamin D content of codliver oil is observed after ultraviolet irradiation, excess exposure being destructive not only of vitamin A but of vitamin D as well

^{**} Hess and Weinstock, Proc Soc Expl. Biol. Med., 23, 407 (1926), Bills J. Biol. Chem., 72, 751 (1927); J. Nutrition, 13, 435 (1937); Notewarp Tids Kjemi, Bergresen Met. No 4, 415 (1937).

^{**}O Nelson, et al. Ind Eng Chem. 22, 1361 (1930); 23, 1066 (1931); Truesdail and Boynton Ind. Eng Chem. 23, 1136 (1931), Brooks, Abernethy, and Vilbrandt J. Am Conard Brocken J. 16, 482 (1922), Copping Brocken J. 28, 1516 (1934), Drummond

and Gunther J Exptl Biol , 11, 203 (1934)

¹⁰² Leigh Clare Biochem, J . 21, 368 (1927)

⁴²³ Belloc, Fabre, and Simmonet. Compt. rend., 191, 160 (1930); Russell Nature, 126, 472 (1930).

⁶⁴ lless, Bills, and Honeywell J. Am Med Assoc. 92, 226 (1929). 193 Hawk Reported before the American Chemical Society, Minneapolis Meeting 1929 Of 1779 hvers examined, those from female cod (58 6 per cent of the total) averaged about 288 ml in size and contained 37 1 per cent oil, while those from the male averaged 163 ml and contained 433 per cent oil

Vitamin D has a very limited distribution among the common foodstuffs Of these, egg yolk probably takes first rank, although this source is quite variable, depending upon the hen's ration and upon the amount of exposure to ultraviolet radiation. The vitamin D content of hutterfat is relatively low, in contrast to its vitamin A content, but is likewise somewhat higher in summer than in winter. Human milk is an even poorer source of the antirachitic vitamin than cow's milk.

Processes used for increasing the antirachitic potency of whole or evaporated cow's milk include: (a) feeding activated yeast or sterols to cows, a small fraction of the vitamin D being transmitted to the milk; (b) ultraviolet irradiation of the milk; (c) fortification of the milk with natural or synthetic vitamin D.** The last is the principal method in use in this country today. Vitamin D milk produced by direct fortification is standardized to contain 400 U.S.P. units per quart (whole milk) and provides adequate safeguard against rickets

Antirachitic activation of foods by ultraviolet irradiation is dependent

upon the presence of the provitamin. Hence certain food products—e.g., milk, egg yolk, yeast—are capable of much greater activation than others, like cornstarch, egg white, or certain cereals. Careful control is required in irradiation processes to avoid the development of rancidity or disagreeable odors, and to guard against destruction of vitamin A, riboflavin, pyridovine, and (by overirradiation) of the vitamin D which is formed.

Plant foods and vegetable oils are notonously deficient in vitamin D. Excellent sources of vitamin A though they are, the green leafy vegetables lack the antirachitic factor in significant amounts even when grown in the summer months. Sun-drying of hay or atfalfa results in a considerable increase in antirachitic potency. Some activation occurs during fermentation of coca, owing to solar radiation of the sterols present in contaminating yeasts or fung.

Certain cereals, notably outmeal, contain a factor which actively inhibits calcification "Not only is part of their phosphorus content nona-similable, but it occurs in the form of ino-itol hexaphosphoric acid or phytin" which binds calcium and iron and thus interferes with their absorption.

spectrophotometric absorption of the cluate at 265 mµ, chemical assay of irradiated sterols or similar high-potency materials is possible. 409 Similar methods employing modified antimony trichloride reagents have heen used for the assay of fish liver oils, but with less success owing to their lower vitamin D potency and the interference of vitamin A. The antimony trichloride reaction differentiates hetween ergosterol and 7-dehydrocholesterol, which are indistinguishable spectrophotometrically. 410

Green has developed a promising chemical procedure for vitamin D in fish liver oils411 involving a series of steps designed to eliminate the interference by sterols and vitamin A with color reactions. The unsaponifiable extract is chromatographed on acid-activated floridin to destroy vitamin A: the carbon tetrachloride cluate is evaporated to dryness and taken up in 72 per cent ethanol to precipitate interfering provitamins and sterols: the filtrate is dried and taken up in chloroform, and the vitamin D ("anparent ealciferol") is estimated spectrophotometrically after reacting with the antimony trichloride-acetyl chloride reagent of Nield, Russell. and Zimmerli.412 While the method is claimed to give good agreement with bioassays, the "detailed steps necessary for the analysis of vitamin A-containing oil have generally to be decided for each particular sample" on the basis of the ratio of vitamin A to vitamin D and the percentage of unsaponifiable matter. By means of an iodine trichloride reaction the method has been adapted for the partial analysis of the irradiation products of ergosterol and 7-dehydrocholesterol, some of which remain unidentified 413

BIOLOGICAL METHODS

The quantitative study of vitamin D activity is hased on the determination of the comparative amounts of test materials and of standard vitamin D preparations necessary to induce healing of rickets in rats or to maintain aormal calcification in chicks The production of experimeatal rickets of a moderate hut uniform degree of severity within a definite period of time is the primary requisite for reproducibility of results. Various methods differ in respect to the composition of the rachitogenic diet, the criteria for the establishment of the presence and degree of rickets, and the hasis for judging the degree of healing. The rat curative assay (U.S.P.) is employed for the evaluation of the vitamin in foods or pharmaceutical products intended for human (or other mammalian) use; hecause of the relative insensitivity of fowl to vitamin D2, the chick method (A.O.A.C.) is employed in the assay of vitamiu D intended for feeding poultry.

Ordinarily diets used for the production of experimental vitamindeficiency diseases are characterized by being complete in all respects (i.e., as regards proteins, calories, minerals, and vitamins) except for

Ewing, Kingsley, Brown, and Emmett Ind Eng Chem, Anal. Ed., 15, 301 (1943);
 Ewing, Powell, Brown, and Emmett Anal Chem, 29, 317 (1948).
 Lamb, Müller, and Beach: Ind. Eng Chem, Anal. Ed., 18, 187 (1946).

⁴¹¹ Green: Brochem J., 49, 243 (1951).

⁴¹¹ Nield, Russell, and Zimmerli J. Biol Chem , 136, 73 (1940).

To prepare this reagent dissolve 18 g antimony trichloride in 100 ml, purified chloroform, filter, and add 2 ml redistilled acetyl chloride. Prepare fresh dails

⁴¹³ Green Biochem, J., 49, 232 (1951).

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Ordinarily diets used for the production of experimental vitamindeficiency diseases are characterized by being complete in all respects the absence of the vitamin in question. This, however, is not so in the case of the vitamin D assay with rats because decalcification does not occur readily in the absence of vitamin D unless the normal calcium; phosphorus ratio is disturbed. The analog of elinical rickets may be produced experimentally in rats by feeding a vitamin-D-free diet possessing a Ca:P ratio of ahout 45:1 instead of the normal ratio ranging from 1:1 to 2.1.





Fig 289 (Left) Active Rickets Roentgenograph of knee joint of rat on Steenbock rachitogenic diet No 2965 for 17 days Note especially broad bands of uncaleified cartilage near ends of tibia and femur.

Fig 290 (Right) Healed Rickers
Roentgenograph of knee joint shown
in Fig 289 after the rat had been fed a
codiner-oil concentrate 14 days

After 18 to 21 days on a standard rachitogeme diet, the animals develop a decalefied zone in the epiphyseal region of the long bones (the so-called "rachitic metaphysis") which may be readily demonstrated roentgenographically (see Fig 289). At his time gross symptoms of rickets, such as shambling gait or enlargement of the knee joint, are not yet apparent Hence it is desirable in order to establish the presence of the rachitic lesions that the animals be x-rayed or fluoroscoped or that some of them be sacrificed and their leg bones examined histologically

After three weeks the bone ash values, expressed on a dry fat-free basis, drop from 35-45 per cent, the value at the time the rats are placed on the rachitogenic diet, to 27-35 per cent. At this age the normal ash content should be about 50 60 per cent. Similarly, the morganic phosphorus content of the blood decreaces from the normal level (for rats) of 8-10 y 100 ml. of serium, to about 2-3 mg. "Uning this period the

animals continue to grow, but at a somewhat subnormal rate, the more rapidly growing animals exhibiting rickets of greater seventy

Following the preparatory period, the animals are fed the same diet alus the material to be assayed, which may be incorporated in the diet or preferably fed as a supplement. Negative controls are allowed to contime on the basal diet without antiraelutic supplements, while positive controls receive a daily dose of a standard source of vitamin D. viz. the International or U.S. Pharmaconeia Vitamin D. Reference Oil It is important that food materials fed as supplements do not appreciably after the Ca P ratio of the diet. In the event of this possibility the assay should be controlled by including the ash of the test material in the diet of the negative controls, or preferably feeding the supplement in the form of its ether extract.

Rats which lose weight or fail to consume sufficient food must be discarded, because starvation is attended by a remobilization of tissue phosphate and spontaneous healing of rickets. The animals must be shielded against exposure to similarly or even against undue exposure to filtered day light

The duration of the test period depends upon the enterior employed for healing. In the rocutgenographic method, a six-day curative period may be used 415 (See Figs 289 and 290) This method has the advantage of permitting each animal to serve as its own control, and may be ean veniently combined with the histological technique as prescribed by the US Pharmaeopeia The examination of stained sections of the leg hones (Line Test, see Fig. 291) is conducted at the end of the assay perrod

The feces of rats on a rachitogenic diet tend to become alkahne, but return to the normal shehtly acid condition during the process of healing 416 The increase in pH is localized in the cecum and colon rather than in the small intestine, which is the main site of calcium and phosphorus absorption A concomitant shift of urinary pH occurs in the opposite direction These pH changes in rickets are reversible with vitamin D therapy 417 Efforts to adapt fecal pH shift to the quantitative estimation of vitamin D418 have fuled owing to the small (about one pH unit) and variable difference and the interfering effect of extraneous factors

Fish liver oils and vitamin D concentrates are important constituents of poultry rations Since the rat assay is an madequate measure of the vitamin D value of certain fish oils and sterols for avian species a quantitative method of vitamin D assay employing chicks has been adopted by the Association of Official Agricultural Chemists 419 This is 3 preventive

[&]quot;Poulsson at 1 Love skold Brochem J 22 135 (1978)

"Zucker and Mattner Proc Soc Expl Bud Ved 21 156 (1971)

"Steenbook Bellin and West J B of Chem 193 843 (1951)

"Steenbook Bellin and West J B of Chem 193 843 (1951)

"Jepl cott and Breharach Bucchem J 29 1351 (1976) 22, 60 (1978) J Bud Chem 80 1751 (1978) Covard Q art J Pharm. 1 2" (1978) (1978) Over J Bud Chem 80 487 (1978) Covard Q art J Pharm. 1 2" (1978) 10 (1978) Over J Bud Chem 19 17 (1978) A 17 (1978) And H Halvord And Lall A. J Live Official Laple and B Hs 1 trition 12. 450 (1986) and H Halvord and Lall A. J Live Official Laple 1978 (1987) and H Halvord 1978 (1987) Green Idd 29 135 (1987) (193") Griem Ibid 20 138 (1937)

technique based on a comparison of the average bone asb content of the assay group with that of reference groups receiving graded doses of the Standard One AOAC chief unit of vitamin D is equal in biological activity for the chiek to one unit of vitamin D in the USP Reference Standard

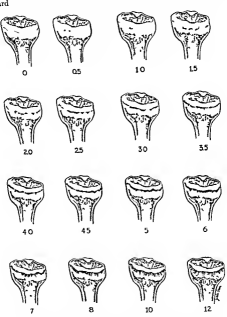


FIG 291 CHART FOR INTERPRETATION OF LINE TEST OF TIBIA

Based on 7-day line test responses according to the U.S. Pharmacopeta procedure. Note that on this scale in numerical ratings 0.5 corresponds to minimal recalification (positive microscopic evidence of calcification). 40 corresponds to a narrow continuous. line of calcification along the metaphy scal junction and 12 falls somewhat alort of complete healing but is the maximal response in a 7-day test.

Whole.

ASSAYS FOR VITAMINS A AND D

Riological Methods of the U.S. Pharmaconeia XIV.430

Departures As used herein, unless the context otherwise indicates, the ferm assayer means the individual immediately responsible for the interpretation of the assay the term assay aroun means the group of rats to which the assay oil shall be administered during the assay period, the term assay oil means the oil under examination for its vitamin potency, the term assay period for the utamin A assay means the interval in the life of a rat between the last day of the depletion period and the twenty-ninth day thereafter or between the last day of the depletion period and the death of the rat, the term assay period for the silamin D assay means, the interval in the life of a rat between the last day of the depletion period and the eighth day thereafter the term assemble means the procedure by which rate are selected and assigned to groups for the nurpose of feeding, care, and observation, the term control group means the group of rats receiving no assay oil during the assay period, the term declining weight means the condition of a rat when its hody weight on any given day is equal to or less than the hody weight of the rat on the seventh day prior to the given day .421 the term depletion period means the interval in the life of a rat between the last day of the preliminary period and the first day of the assay period, the term dose means the quantity of the USP Vitamin A Reference Standard or of the assay oil to be fed at specified intervals to a rat during the assay period, the term fed means made readily available to the rat or administered to the rat liv mouth, the term ground cluten means the clean, sound product made from wheat flour by the almost complete removal of starch, and contains not more than 10 per cent of moisture, and, calculated on the water-free hasis, not less than 14.2 per cent of nitrogen, not less than 15 per cent of nitrogen free extract (using the protein factor 57), and not more than 55 per cent of starch (as determined by the diastase method (*2), the term group for the ulamin A assay (higheren)) means six or more rats maintained on the same required dietars regimen during the assay period, the term group for the mamin D assay means seven or more rats maintained on the same required dietary regimen during the assay period. the term ophthalma means a pathological state of the eve and/or the conjunctiva and/or the tissues anatomically related to the eye, readily discernible macroscopically and usually associated with vitamin A deficiency, the term preliminary period means the interval in the life of a rat between the seventh day after birth and the first day of the depletion period, the term rachitogenic diet means n uniform mixture of the food materials, and in the proportions named, in either of the following formulas

Rachitogenic Diet No 1	Rachitogenic Diet No 2
Yellow Maize, ground 33 per cent	Whole Yellow Maize, ground 76 per cent
777 / 1 00 /	0 101

Whole Wheat, ground 33 per cent Ground Gluten 20 per cent 15 per cent Calcium Carbonate (USP) Ground Gluten 3 per cent Gelatin 15 per cent Sodium Chloride (U.S.P.) 1 per cent Calcium Carbonate (USP) 3 per cent

Sodium Chloride (USP) 1 per cent

433 Association of Official Agricultural Chemists Official and Tentative Vethods of Analysus 6th ed 1945 p 410 method II, p 577, method 34 53

⁴²⁰ Grateful acknowledgment for permission to reproduce these methods is made to Dr Lloyd C Miller and the Board of Trustees of the U.S. Pharmacopeial Convention

⁴²¹ In this connection it is well to remember that rats manifest a diurnal variation in body weight averaging about 2 g. Because of the severity of avitaminosis and the high mortality that often results when rats remain too long on the depletion diet many workers prefer to reduce the period of declining weight' to four or five days Differences in pathological symptoms which arise at the end of the depletion period are responsible for much of the variation encountered in hiological assays for vitamin A. It has therefore been suggested that preventive rather than curative technique might yield more uniform results

The term estamin A test diet means a food material consisting of the following proportions of the named ingredients of the quality specified

Vitamin A Test Diet

18 per cent
4 per cent
8 per cent
65 per cent
5 per cen

No less than 3 USP Units of vitamin D shall be provided in each gram of diet and this vitamin shall be carried by the yeast or the cottonseed oil. The ingredients of the vitamin A test diet shall be free from vitamin A or shall have been treated so as to reduce the vitamin A content to such a degree that when the vitamin A test diet is fed to the control group two-thirds or more of the rats shall manifest, prior to the eleventh day of the assay period symptoms of vitamin A deficiency characterized by both declining weight and ophthalmia

Salt Mertura

Sait Vixture	
Calcum Carbonate (USP)	68 6 g
Calcium Citrate (USP Reagent)	308 3 g
Calcium Biphosphate (USP Reagent)	112 8 g
Magnesium Carbonate (USP)	35 2 g
Magnesium Sulfate, Anhydrous (USP Reagent)	38 3 g
Potassium Chloride (USP)	124 7 g
Dibasic Potassium Phosphate (USF Reagent)	218 8 g
Sodium Chloride (USP)	77 1 g
Cupric Sulfate (U S P)	0 48 g \
Ferric Ammonium Citrate (USP)	96 71 g
Manganese Sulfate (USP Reagent)	1 24 g
Ammonium Alum (USP Reagent)	0 57 g \ 16 2 g
Potassium Iodide (USP)	0 25 g (
Sodium Fluoride (USP Reagent)	0 75 g
To make	100 00 g
To make	1000 0 g

Mix the finely powdered salts uniformly

The biological assay of an oil for vitamin A and vitamin D potency shall be by comparison with the USP Vitamin A Reference Standard 42 and USP Vitamin D Reference Standard respectively, by assay procedures conforming in all respects to the following specifications

Assay for Vitamin A-Biological Method

For products for which the spectrophotometric method of vitamin A assay is not applicable use the biological method which follows

The Vitamin A biological assay comprising the recording of observations of groups of rats throughout specified periods of their lives will be being maintained on specified dietary regimens, and the interpretation of such data is as follows:

⁶³ The USP Vitamin A Reference Standard consists of an encapsulated solution of crystalline vitamin A acetate in cotton-seed oil

This of solution is assigned a value of 10 000 USP vitamin A units per gram The content of each capsule is assigned a value of 2500 USP vitamin A units

PRELIMINARY PERIOR Throughout the preliminary period each rat shall be raised under the immediate super isson of, or according to directions specified hi, the assaver Throughout the preliminary period the rats shall be maintained on a dietary regimen which shall provide for normal development in all respects, except that the supply of vitamin A, or precursors of vitamin A, shall be limited to such a degree that rats weighing between 40 and 50 g and not exceeding 28 days of age and subsisting on a suitable vitamin A deficient ration and water for an interval not exceeding 45 days shall manifest symptoms characteristic of vitamin A deficients.

DEPLETION PERIOD A rat shall be suitable for the depletion period when the age of the rat does not exceed 28 days, and if the body weight of the rit shall exceed 39 g, and does not exceed 50 g, and if the animal manifests no evidence of injury, or disease, or anatomical abnormality which might hinder growth and development. Throughout the depletion period each rat shall be provided with the vitamin A test did not not not the rate of the provided with the vitamin A test did not not the shall be available to the animal

ASSEMBLING RATS INTO GROUPS FOR THE ASSAY PERIOD Rats which are soutable for the assay period shall be assembled into groups. For each assay oil there shall be one or more assay groups. In the assay of one assay oil there shall be provided at least one control group and at least one reference group, but one control group and one reference group may be used for the concurrent assay of more than one assay oil The interval of assembling rats into groups shall not exceed 60 days but the depletion period of rats in any one litter shall not vary by more than 14 days. On any one day during the interval of assembling rats into groups, the total number of rats that shall have been assigned to make up any one group shall not exceed by more than two the number of rats that shall have been assigned to make up any other group When the assembling of all groups shall have been completed, the total number of rats in each group shall be the same, and the number of rats of one sex in each group shall be the same oot more than three rats from one litter shall be assigned to one group. When the assembling of all groups shall have been completed, the average weight of the rats in any one group on the day beginning the assay period shall not exceed by more than 10 g the average weight of the rats in any other group on the day beginning the assay period

Assay Person A rat shall be suitable for the assay period, provided that the depletion period shall have exceeded 18 days and shall not have exceeded 45 days. and provided that a rat shall manifest evidence of vitamin A deficiency characterized by declining weight and/or ophthalmia. Throughout the assay period each rat of the control, reference, and assay groups shall be kept in an individual cage and shall be provided with the vitamin A test diet and water (USP), ad libitum Throughout the assay period each rat in any assay group shall be fed two doses each week of the assay oil, and throughout the assay period each rat in any one reference group shall be fed two doses each week of the USP Vitamin A Reference Standard The first dose of the assay oil or of the U.S.P. Vitamin A Reference Standard shall be fed on the day beginning the assay period, and succeeding doses shall be fed at intervals of not less than 3 nor more than 4 days The USP Vitamin A Reference Standard and/or the assay oil may be diluted before feeding with cottonseed oil (USP) Diluted oil shall be stored in the dark at a temperature not exceeding 10° C. The period of storage sbull not exceed 7 days Not more than 0 2 ml of the diluted oil shall be fed in a single dose During the assay period all conditions of environment shall be maintained as uniformly as possible with respect to the assay, reference, and control groups

RECORDING OF DATA On the day beginning the depletion period and at intervals of not more than T days for the first 21 days of that period, there shall be a record made of the body weight of each rat. From the twenty first day of the depletion period until the end of the assay period a record shall be made of the body weight and eye condition of each rat at intervals not exceeding 5 days. The eye condition shall be

sections of the bone shall be mised in distilled water and shall then numediately be immersed in a 2 per cent aqueous solution of either nitrate for 1 minute The sections shall then be rinsed in distilled water and the sectioned surfaces of the bone shall be exposed in water to daylight or other source of actinic light until the calcified areas have developed a clearly defined stain without marked discoloration of the uncalcified areas

Records shall be made immediately of the extent and degree of calcification of the rachitic metaphysis of every section. It shall be permissible to use modifications of the described procedure for staining provided that such modified procedures clearly differentiate between calcified and uncalcified areas.

RECORDING OF DATA On the day beginning the assay period and on the seventh day thereafter, a record shall be made of the body weight of each rat A record shall be made of the quantity of relutogenic diet consumed per rat during the assay period Numerical values shall be assigned to the extent and degree of catcification of the raclutic metaphysis of the bones examined by the line test so that it will be possible to average the performance of each group A Line Test Chart, that may be used for this purpose is presented below ***

VITAMIN D POTENCY OF THE ASSAY OIL In determining the vitamin D potency of the assay od the average performance of groups with respect to healing of the rachitic metaphy as shall be considered provided that the average performance of a reference group with respect to calcification of the rachitic metaphysis shall be determined by the data from rats which individually show an extent and degree of calcification in the rachitic metaphysis as determined by the line test equal to or greater than a condition described as a positive macroscopic evidence of calcification, but no greater than an extent and degree of calcification represented by the value 8 in the Line Test Charters presented below The data from a reference group shall be considered valid for establishing the vitamin D potency of the assay oil only when two-thirds or more, but not less than seven rats show individually an extent and degree of calcification of the rachitic metaphysis equal to or greater than a condition described as positive macroscopic evidence of calcification but no greater than an extent and degree of calcification represented by the value 8 in the Line Test Charters presented below. The data from an assay group shall be considered valid for establishing the vitamin D potency of an assay oil only when two-thirds or more, but not less than seven rats, show in dividually an extent and degree of calcification of the rachitic metaphysis equal to or greater than a condition described as positive macroscopic evidence of ealcification The data from a rat shall be considered valid for establishing the average performance of a group only on the condition that the weight of the rat at the termination of the assay period shall equal or exceed the weight of the rat on the beginning day of the assay period and that the rat has consumed 28 g or more of the rachitogenic diet dur ing the assay period and on the condition that the rat has consumed each prescribed dow of assay oil within 24 hours from the time it was fed

CALCULATION OF VITAMIN D POTENCY

Let R equal the total quantity of USP Vitamin D Reference Standard in milligrams fed to each rat in a reference group and necessary to produce in a reference group an average extent and degree of calcuffeation C not less than a condition described as positive macroscopic evidence of calcuffeation but less than the degree of calcuffeation represented by the value 8 in the Line Test Chart

Let I equal the total quantity of assay of in milligrams fed to each rat in an assay group and necessary to produce in an assay group an average extent and degree of calcification equal to or greater than C

 $^{^{\}rm 418}$ This chart of the line test responses as seen in the radius has been omitted. See footnote 424 $\,p\,$ 1265.

MINIMUM STANDARD If the product of $\binom{R}{4}$ × (units per g of vitamin D contained in the USP Vitamin D Reference Standard) is equal to or greater than the minimum standard in USP units per g for the oil assayed, then the assay oil meets the minimum standard for vitamin D notence ***

Maximum Standard If the product of $\binom{R}{A}$ × (units per g of vitamin D contained in the USP Vitamin D Reference Standard) is equal to or less than the maximum standard in USP units per g for the oil assayed, then the assay oil meets the maximum standard for vitamin D potency ⁴²⁴

VITAMIN E (TOCOPHEROLS)

The essentiality of vitamin E for optimum nutrition has been demonstrated in more than fifteen vertebrate species by restricting the dietary intake of this vitamin and noting deficiency symptoms. The discovery of vitamin E in 1922 was based upon experiments of H M Evans, of the University of California, showing that for normal reproductive function rats required a new fat-soluble factor present in crude vegetable oils and subsequently identified as isomers of tocopherol Later studies showed that this new vitamin was also essential for normal muscle metabolism. to maintain the integrity of the central nervous system and the vascular system, and for a variety of other physiological functions. The pathology of the symptoms of vitamin E deficiency has been described in great detail, particularly by Pappenheimer and Goettsch and by Mason However, the exact brochemical mechanism wherehy vitamin I functions in diverse metabolic processes in the body is still unknown. This and the direct experimental production and recognition of vitamin E deficiency in human subjects constitute two of the most important problems in the field of vitamin E research Estimates of the human requirements for this vitamin are hased upon indirect evidence, such as the mean daily per capita consumption of vitamin E and the similarity of tissue pathology seen in premature infants and adults suffering from faulty fat absorp tion to that seen in vitamin E deficient animals

Physiological and Clinical Aspects of Vitamin E Not all of the tocopherol in foods is completely available when ingested The α form is more efficiently absorbed than the non α tocopherols. Tocopherols in cereal grains leafy vegetables, and practically any natural foodstiff are bound in such a way that during passage through the digestive tract some of the tocopherol is evereted in the feces. Balance studies in normal human heings above as much as 60 per cent of ingested α tocopherol appearing in the feces. Conditions interfering with fat absorption (bihary diseases, pancreatic insufficiency, innerial oil ingestion etc.) further reduce the amount of this vitamin absorbed. No free or conjugated tocopherol is exerted in the urine. Both tocopherol and tocopherol esters when ingested appear in the blinod and are transported as the free form

 α Tocopherol is deposited in all tissues. The pituitary and adrenal glands have especially high concentrations of tocopherol, over 30 mg per

 $^{^{639}}$ The U.S.P. Unit of vitamin D is defined as the activity of 0.0°5 μg of vitamin D, contained in the U.S.P. Vitamin D Reference Standard

cent, but the bulk of the body stores of tocopherol is in the adipose tissues and muscles

Vitamin E deficiency symptoms develop in different tissues and at different rates depending upon species. The rabbit and lamb show muscular lessons and paralysis within a few weeks on a tocopherol-free det. The rat develops testicular degeneration and the chick shows softening of the brain (encephalomalacia) within six weeks. However, the lamb, rabbit, and herbivores in general eventually develop reproductive dysfunction, the rat shows muscle dystrophy and paralysis in about 12 to 18 months, and the chicken eventually develops both muscular and reproductive abnormalities. No common denominator of physiological function of a tocopherol or of reasons for species specificity and tissue susceptibility has been established, this makes it necessary to consider separately the physiology of individual tissues and organs.

Complex biochemical changes in the muscle tissue in chronic vitamin E deficiency are followed by histological lesions characteristic of muscular dystrophy. The muscle shows either a gradual necrosis and replacement of musclo fihers with fat and connective tissue, or an explosive form of dystrophy involving larger portions of the muscle and accompanied by edema, leukocytic infiltration, and fragmentation of muscle fibers The former is typified by the muscular paralysis which develops in the adult hamster and has its histological counterpart in humans in progressive muscular dystrophy and dermatomyositis The explosivo type of syndrome is exemplified by paralysis in suckling rats late in factation and by muscle dystrophy in guinea pigs and rabbits. Heart musclo is affected like skeletal musele by vitamin E deficiency and rats, rabbits, sheep, and cows have all shown typical lesions of degeneration, pigment deposition, electrocardiogram changes, and functional failure Vascular abnormalities develop during the production of vitamin E deficiency in all species studied The dog in particular, when subjected to a high fat diet combined with kidney injury, develops a reversible arteritis resembling that seen in human arteriosclerosis

Demyelimization, ghosis, and distortion of the axon pattern in the spinal cord are seen in vitamin E deficient rats, giving rise to hypalgesia

and progressive paresis

The vitamin E deficient male animal shows testicular atrophy and sterility due to irreversible degeneration of the semiinferous tubules and to failure of spermatogenesis. In the vitamin E deficient female, a reversible type of reproductive dysfunction occurs. Ovarian function is normal but interine physiology is disturbed. There is partial failure of implantation, and those fertilized on a successfully implanted grow and develop only to a certain stage at which time the fetuses show generalized hemorrhage the and are aborted or resorbed. Administration of a-tocopherol during the first half of gestation permits normal fetal divelopment and partirition. This reversibility of reproductive function in the female rat serves as the basis for the bioassay method most generally used to evaluate vitamin F potency.

The normal resistance of red blood cells to rupture, both in viro and in ritro, is markedly reduced in vitamin E deficiency. Administration of

 α -tocopherol to the patient or addition of α -tocopherol to the *in vitro* system returns the resistance of the **erytbrocytes** to normal. This improvement in function of red blood cells by α -tocopherol serves as the basis for a bioassay for vitamin E activity.

The finding that vitamin E metabolism plays an essential role in cardiovascular pathology in lower animals bas resulted in attempts to employ tocopherols therapeutically. Cases of angina pectoris, coronary insufficiency, peripheral vascular diseases, etc. have been reported by some investigators to improve under α -tocopherol therapy, but others have failed to observe any benefit in similar clinical trials

Fatal massive liver necrosis occurs in animals maintained on diets low in vitamin E and sulfur-containing amino acids. Under conditions in which the intestinal flora of the animal are eliminated (germ-free environment, antibiotic treatment), liver necrosis does not develop but the animal dies from lung hemorrhage. α -Tocopherol administration prevents both the liver necrosis and the lung hemorrhage. Possibly this prevention of liver injury by α -tocopherol is merely an example of a physiologic role of α -tocopherol whereby environmental, dietary, and drug stresses are combatted.

Some evidence, both experimental and clinical, points to a sparing effect of α -tocopherol upon hormone action, particularly insulm and the sex hormones This is considered a result of α -tocopherol everting an antioxidant or sparing action upon the hormone metabolism. Various patbological states in poultry and livestock have been shown to be aspects of vitamin E deficiency and to respond to administration of tocopherols Among them are "crazy-chick disease," "cularged bock disease" of turkey poults, "white muscle disease" of calves and colts, "stiff lamb disease," and steatitis or "veillow fat diseases" of mink.

REQUIRENENT. The need for vitamin E has been established qualitatively for a variety of animal species. However, quantitative requirements, either minimal or optimal, for various animals including man, have not been determined.

Diets of good quality consumed by healthy persons in good economic circumstances average between 10 and 25 mg. a-tocopherol daily. Reducing diets and some therapeutic-type diets furnish as little as 4 mg. per day. From a curve relating vitamin E requirements for lahoratory animals to (body weight)⁹, the value for a 70-kg, animal may be extrapolated An estimated requirement for humans of 30 mg. p-a-tocopherol is thus derived. From a consideration of the results of these two approaches, the human requirement may be estimated as 15-25 mg. a-tocopherol daily.

Biopotency. a-Tocopherol and various of its derivatives have different relative potencies depending upon the procedure used for bioassay. Esterified a-tocopherol is more potent by the rat bioassay (in which suboptimal amounts are fed) than an equivalent quantity of free a-tocopherol, probably because of the relative instability of the latter in the intestinal tract (see Table, p 1270). Tests on buman infants and adults with therapeutic doses fail to show this superiority of the ester form. Hence the National Formulary X considers 1 mg of synthetic a-tocopheryl acetate

RELATIVE BIOPOTENCY OF VARIOUS FORMS OF VITAMIN E

Compound	Wolecular WeigH	Relative Biopolency*	International Units per Mg	
N atural		401	0.00	
p–α Tocopherol	430 7	101	0 92	
p-α Tocopheryl acetate	472 8	136	1 36	
p-a Tocopheryl acid succinate	530 8	136	1 21	
p-a Tocopheryl disodium phoephate	554 7	13	0 10	
Synthetic	i	1	ĺ	
racemic-a Tocopherol	430 7	75	0 68	
-a Tocopheryl acetate	472 8	100	1 00 (by definitio	
(International Standard)	1	1		

^{*} Based on feed ng equivalent amounts of a tocopherol

equal to 10 I U and 1 mg of natural α to copheryl acetate equal to 136 I U The free α to copherols on the bass of their molecular weight, have a biopotency of 11 I U per mg for the synthetic and of 149 I U per mg for the natural form

Physiological Antioxinant Activity The fact that protoplasm is an aqueous-lipide emulsion in which metabolic reactions occur at interfaces immediately poses the problem of properly stabilizing the lipide phase a Tocopherol is the only naturally occurring, fat-soluble food antioudant which accompanies lipide through the intestinal tract and during its transport via blood or lymph to storage depots or metabolic sites Other antioudants are eliminated during absorption or in transit, and a tocopherol emerges as the only physiological fat soluble antioudant

The amount of α -toeopherol deposited in the tissues is proportional to the quantity ingested. In certain species—for example, turkeys and pugs—the concentration of α toeopherol in tissue lipides is so low that turkey carcass fat and lard oxidize and become raised even under conditions of low temperature storage. The convexeretes a proportional amount of ingested or stored α toeopherol into her milk and whenever the level fall's belo v. 20–30 μ g per g milk fat the milk has poor stability and off flav or develops.

Chemistry of the Tocopherols The four compounds in vegetable oils possessing vitamin E activity have been siolated and their structures proved They were named locopherols (Greek, "young bearing"), and found to differ only in the extent of methylation of the ebroman nucleus α Tocopherol was sinthesized by L. I. Smith and by Karrer, independently in 1938. The structure of α tocopherol is shown on p. 1271 with those carbon atoms marked where asymmetry may occur, thus giving rise to isomers. Natural α tocopherol is the p form whereas synthetic vitamin Γ consists of a mixture of isomers usually designated as racemie place tocopherol.

The other tocopherols β -, γ -, and δ -tocopherols, have been synthesized and tharseterized However they are considered to be less important, since they are less effective physiologically than a tocopherol. The structural differences of the various tocopherols are shown in the follow-

a Tocopherol

* Points of asymmetry Natural-occurring α tocopilerol is the p form whereas synthetic α tocopherol is a mixture of isomers

mg scheme The β form has about one-third the biopotency of α -tocopherol, the γ form about one-twelfth, and the δ form about one hundredth

The Tocopherol Nucleus

Name

Substituents at Positions

	5	~	8
α Tocopherol = Trimethyl derivative β ' = Dimethyl " γ ' = Dimethyl " δ- " = Methyl '	CH.— CH.— H— H—	CH ₁ — CH ₂ — H—	CH.— CH.— CH.—

α-Tocopherol is easily destroyed by oxidation Esterification with acetic, succinic phosphoric, and other acids is commonly practiced to stabilize the vitimin These esters have essentially the same potency as α-tocopherol except that under certain test conditions they bioassay higher than the free tocopherol hecause of their enhanced stability

Other changes in the molecule, such as alteration of the side chain, etherification at position 6, or substitution of the ring methyl groups, markedly reduce or completely eliminate vitamin potency Oxidation of a-tocopherol, which progresses through an epocide intermediate, a hydroquinone, and a free radical semiquinone to tocopherol quinone, destroys the vitamin E potency for most animal species However, the epocide intermediate is physiologically active for the rat, the tocopherol hydroquinone will prevent or cure muscle degeneration in the hamster and the final oxidation product, tocopherol quinone, bas vitamin E activity for the rabbit

estimation of tocopherol with ferric chloride dipyridyl reagent measure total tocopherois, and some procedure for obtaining only the a-tocophero content is required in foods where the tocopherol concentration is low y- + 5-tocopherol concentration is measured by diazotization with dia nisidine 430 (8-Tocopherol is not measured, but this form occurs only in wheat products) The a-tocopherol content is obtained by subtracting the combined value for 7- + 8-tocopherols from the total tocopherol value

In preparations where at least 0.5 mg of β - + γ - + δ -tocopherols are available, analysis of their total content is preferably done by the nitroso method 431 This value for the combined amount of all non-a-tocopherols, subtracted from the total tocopherol value determined by the ferric chloride dipyridyl method, has been used to give a measure of the a-tocopherol content

Separation, isolation, and direct measurement of a-tocopherol in extracts of biological materials has been done semiquantitatively by means of paper chromatography. Brown 42 has reported separation of the tocopherols, using petrolatum coated paper strips and 75 per cent ethanol as solvent, and has estimated the Individual tocopherol content of various animal and vegetable f1831168

BIOLOGICAL METHODS

Erans Resorption-Gestation Method, Modified by Moson and Harris. 411 This bigassay is based upon the ability of a-tocopherol, administered to vitamin E deficient female rats during the first half of pregnancy, to prevent fetal death and resorption Female rats with low vitamin E reserves are raised on the following vitamin E low diet

Casein, commercial unextracted	20	per	cent
Carbohydrate, starch, or sucrose	56		**
Fat, lard, or olive oil	10		46
Salt mixture	4		**
Brewer's yeast, dried	10		**
Vitamins A and D, added to fat component			
to supply 10 units vitamin A and 1 unit			

When the females are 150 g in weight, they are mated with normal males Vaginal smears are made daily until sperm (sign of positive mating) are observed. The pregnant females are placed in individual cages and assigned at random to control or test groups

Groups of six to ten rats are set up on four or five levels of both the unknown material being assayed and the standard (The International Vitamin E Standard is a preparation of DL-a-tocopheryl acetate in olive oil. The bio logical activity of 1 mg of this DL α -tocopheryl acetate = 1 International Unit of vitamin E.)

The vitamin E supplement is fed orally in five daily doses prior to the tenth day of pregnancy On the twentieth day of pregnancy the rats are killed and autopsied The uterus is inspected for live, dead, and resorbed fetuses Only those animals with four or more implantation sites are considered A "positive" response is recorded for animals with one or more viable fetuses, and a

vitamin D per gram of diet

¹²⁰ Quarle J Am Chem Soc 66 308 (1944) 121 Quarle J Biol Chem 175 600 (1948)

¹³ Brown Biochem J 52 523 (1952)
14 Mason and Harris Biol Symposia 12 459 (1947)

"negative" for those with no living fetus. A dose:response curve is constructed relating dose (logarithm) to response (litter efficiency, i.e., percentage of positive responses expressed as probits "1) for both the standard and the unknown. These are compared at the 50 per cent end point (the Median Fertility Doses) and the potency of the unknown is expressed in terms of international Units of vitamin E.

Rose-Gjörgy Erythrocyte Hemolysis Test.*** Adult rats, two to three months old, have red blood cells which are normally resistant to the bemolytic action of dialuric acid, hydrogen peroxide, and similar reagents. Changing the animals to a vitamin E low diet for even a few days causes the red cells to become susceptible to hemolysis. Administering various levels of a-tocopherol, either prophylactically or curatively, to the rats during the depleting period and measuring the degree of red cell hemolysis under arbitrary stradard conditions vives a satisfactory bloassay procedure.

Procedure. Rats receiving various levels of pure α -tocopherol as standard and other groups receiving similar levels of tocopherol in the unknown material, both within the range to furnish 0.1 to 0.4 mg. α -tocopherol per day, are used. The supplements are fed daily for two weeks and on the fifteenth day blood samples are takeo and tested. Obtain blood from tall of rats, 10 to 12 drops in a graduated 15-ml. ceotifluge tube containing 2 ml. anticoagulant (equal parts of 0.9 per cent NaCl and 1 per cent sodium citrate solutions). Centrifuge and discard supernatant fluid. Dilute the red cells to a 5 per cent suspeesboo with 0.9 per cent NaCl solutioo.

Pipet 0.25 ml. erythrocyte suspension into each of three test tubes or colorimeter tubes. To one (control) add 4.75 ml. of salioe-buffer solution; of to tube 2 add 4.75 ml. H₂O, which hemolyzes red cells completely; to tube 3 add 4.45 ml. saline-buffer solution; and 0.30 ml. dialuric acid solution, and Mix and incubate for 30 minutes at 37° C. Allow to stand at room temperature for at least two hours, invert tube, cootrifuge, and measure in the colorimeter the red color from the hemolyzed cells.

CALCULATION The reading of the control sample is subtracted from readings of both the test sample and the completely hemolyzed sample. The percentage of hemolysis is calculated as the ratio of the corrected test-sample reading ×100, to the corrected hemolyzed-sample reading.

Human infants, especially premature, have red blood cells which are hemolyzed by dilute hydrogen perovide Administration of α -tocopherol, either in vivo or in vitro, prevents this hemolysis ⁴³⁵

Other bioassays have been developed and used to a limited extent For example, young male rats raised for three months or more on vitamin E deficient diets supplemented with various levels of a-tocopherol, 0 0 to 1.0 mg. daily, show testis weight proportional to the dose of a-tocopherol.

⁴⁴ Converting percentage responses to probate (a mathematical convention), by reference to tables, results in a straight-line dose response curve instead of a sigmoid-shaped curve. This makes it easier to derive a median value from the data (Bliss. Ann. Applied. Blod., 22, 134, 1935).

⁴³⁵ Rose and Gyorgy Proc Soc Expl Biol Med , 74, 411 (1950)

⁴²⁸ Saline Phosphate Buffer, pH 74 02 M KH1PO4, 50 ml; 02 M NaOII, 39 34 ml; and water up to 200 ml Mix with equal volume of 09 per cent NaCl

W Dialuric Acid Solution 1 mg/ml saline phosphate buffer solution (equal parts of phosphate buffer and 0 9 per cent NaCl solution) Since dialuric acid is easily oxidized, the solution should be prepared immediately before use

⁴¹¹ Gyorgy, Cogan, and Rose, Proc Soc Exptl Biol Med. 81, 536 (1952).

"negative" for those with no living fetus. A dose: response curve is constructed relating dose (logarithm) to response (litter efficiency, i.e., percentage of positive responses expressed as prohits (34) for both the standard and the unknown. These are compared at the 50 per cent end point (the Medlan Fertility Doses) and the potency of the unknown is expressed in terms of International Units of vitamin E.

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versely proportional to the dose of tocopherol Rabbits fed a vitamin E deficient dict develop muscle dystrophy in about three weeks which is characterized by creatinuma In the carly and reversible stages of the dystrophy the creating exerction can be reduced temporarily to normal by single doses of vitamin E. The time clapsed from dosing to the return of the ercatinuma to its iintial high level is proportional to the dose of tocopherol

VITAMIN K

In the late 1920's, Dam 429 observed a hemorrhagic syndrome in chicks raised on a diet low in lipides which failed to respond to rich sources of the known vitamins The disease is associated with a reduction in clotting power of the blood, hemorrhages occurring especially in regions exposed to trauma In 1935, Dam advanced the claim that this syndrome was due to deficiency of a now fat-soluble vitamin in green leaves, which he called vitamin K (for Koagulations) Almquist and Stokstad 110 independently announced a similar claim for a factor present in putrefied fish meal. In 1939, the pure vitamin was isolated from alfalfa by Dam and others and from both alfalfa and putrefied fish by MacCorquodale, Binkley, McKec, Thayer, and Doisy 442 Physical and chemical differences in the vitamins thus isolated led to the designation K1 for the vitamin from green leaves and K: for that formed by bacterial putrefaction Investigation of the vitamin K molecule has revealed the presence of a quinoid ring structure, 2-methyl-1,4 naphthoquinone, which alone or substituted in various ways possesses all or part of the physiological activity of the vitamin, thus vitamin K has various natural and synthetic vitamers

Physiological and Clinical Aspects of Vitamin K. The specific type of hemorrhagie diathesis observed in vitamin K deficiency is due to a lowering of the prothrombin level of the plasma (For the role of prothrombin in the coagulation of blood, see p 477) Hypoprothrombinemia may, however, result from other causes such as cirrhosis, chloroform poisoning or Banti's disease, where liver tissue is damaged Vitamin K is without value in hemoplisha, a condition due to a deficiency of thromboplastin and hence a failure of conversion of prothrombin to thrombin Vitamin K does not alter the clotting time of hypoprothrombinemic blood when added in vitro, its function in the body is to facilitate the synthesis of prothrombin in the liver but it has not been definitely established whether the role of vitamin K is that of a precursor or a catalyst Prothrombin synthesis occurs in the hver and a vitamin K "tolerance test" for normal liver function has been proposed. In cases of prothrombin deficiency administration of vitamin K causes the prolonged prothrombin time to return to normal provided the impairment of liver function has

Dam Buchem Z 215 474 (1224) 220 1.08 (1930) Buchem J 29 1273 (1935)
 Umquut and Stokstad J Buc Chem 111 10 (1973) J Aufrition 12, 329 (1936)
 Dam Gegre Glavand Karrer Karrer Rotherlând and Salou on Helt Chim Acta

of MacCor modale Binkley McKee Thayer and Doisy Proc Soc Expil Biol Med 40 482 (1933)

not reached an advanced state 443 The prolonged clotting time associated with hypervitaminosis A responds to vitamin K administration 444

Chicks, ducklings, and other birds fed a vitamin K deficient diet develop severe bemorrhagic lesions both internally and subcutaueously, especially under the wings, on the legs, hreast, indomcu, and neck, and in the intestinal tract. The concomitant anemia terminates in death. In the rat, hacterial synthesis of vitamin K₂ in the intestine, especially when coprophagy is possible, usually obscures the effect of a vitamin-K-free diet.

Simple dietary deficiency of vitamin K is rare, not merely because of the presence of the vitamin in foods, but hecause intestinal microorganisms, especially of the coli group, synthesize the vitamin, which is released upon putrefactive disintegration of the hacterial cells. K avitaminosis in humans is more often associated with impaired absorption of fat (and concomitantly of vitamin K) such as is seen in obstructive jauudice and biliary fistulas, as well as in intestinal disorders like celiae disease and ulcerative cohtis. In these conditions, or after removal of biliary obstrue tions, vitamin K is administered parenterally or orally to prevent or relieve the bleeding tendency. When the flow of bile is obstructed, oral dosago of vitamin K must be accompanied by bile salts to insure its absorption, this may be avoided by the use of water soluble derivatives of the vitamin Vitamin K preparations are also used to treat bemorrhagie emergencies (hypoprothrombinemia) resulting from the use of antiprothrombinemie drugs in thromboembolie conditions (e.g., corouary artery thromhosis)

The most common form of vitamin K deficiency is seen in newborn infants, so-called hemorrhagica neonatorum. The blood protbrombin level is low at hirth, it decreases further during the first few days, rises sharply and then gradually until a normal level is reached between one and two months of age. Whether the initial hypoprothromhinemia is due to poor placental transfer bas not heen established. It is significant that milk, both human and cow, is a very poor source of vitamin K. The rise in protbromhin after hirth is attributed to hacterial synthesis in the intestine Administration of vitamin K to the mother before parturition results in higher hlood protbromhin levels at hirth. Direct dosage of the infant with as little as one microgram of the vitamin is also effective.

The minimum duly requirement of vitamin K has not been established. The extremely low prophylactic dose for the infant, and subsequent synchesis in the intestine, indicate that the dietary requirement, if any, is small. One to 2 mg of the vitamin is capable of correcting most deficiencies, although the therapeutic dose may vary with the severity of the hepatic or intestinal condition. For further discussion of climical aspects, see p. 1293.

Storage and Distribution of Vitamin K. With the possible exception of the liver, the organs and tissues of the hody do not store vitamin K to any significant degree In the hen, even when the ration is rich in this

⁴⁴ Unger Weiner and SI apiro Am. J Clin Pathol 18 835 (1948)
4 Walker E.Jeiburg and Moore Biochem J 41 575 (1947) Quick and Stefanini J Biol Chem 175 945 (1948)

vitamin, only small concentrations are found in the various organs Vitamin K is not found in the unne, the high concentration in the feces is due largely to intestinal synthesis

Occurrence of Vitamin K. The principal natural source of vitamin K is the green leaf or other chlorophyll-containing portions of plants, this form of the vitamin (K₁) has therefore been named phylloquinone. The vitamin is present in high concentration in spinach and alfalfa, while cabbage, cauliflower, seaweed, and carrot tops are good sources. Most seeds, fruits, and roots (including cereals, beans, potatocs, peas) contain little if any vitamin K, although soybean oil, tomatoes, orange peel, and hemp seed are good sources. Milk and eggs are poor sources of vitamin K even when the ration of the cow or hen contains a high level of the vitamin

Chemistry of Vitamin K. Recognition of the quinoid structure of vitamins K₁ and K₂ was due to the work of McKee, et al *** at Washing-ton University, St Louis, and of Karrer and Geiger *** The former group established that these vitamins were derivatives of 2-methyl-1,4-naphthoquinone with substituent groups in the 3 position, K₁ containing the nibvtly group and K₁ a similar but longer (difarnesy) side chain

Vitamin K1 (C11H4O1)
2-methyl-3-phytyl-1,4-naphthoquinone

Vrtamin K₂ (C₁₁H₁₄O₂)
2-methyl-3 difarnesyl-1,4-naphthoguinone

Vitamin K₁ is a light yellow, vised oil, whereas K₁ is a yellow crystalline solid (inclining point 54°). Both are soluble in oil and various fat solventy. The vitamins are sensitive to light and, as would be expected from their quinoid structure, are destroyed by saponification. Vitamins K₁ and K₂ are characterized by ultravolet absorption maxima at wavelengths of 243, 248, 261, 270, and 328 m_µ. The quinoid ring common to these compounds is responsible for their physiological activity. This is shown by the fact that the synthetic compound 2-methyl-1,4-naphthoquinnone, to

⁽¹⁹³³⁾ Was Nac Corquidde Thaser and Doisy J Am Chem Soc., 61, 1295

⁴⁴ Karrer and Geiger Hels Chim Acta 22, 945 (1933)

which the name menadione has been given, is equally as active on a molar basis as the natural vitamers. Menadione has a molecular weight of 172,

Menadione (C₁₁H₈O₂) 2-methyl-1,4-naphthoquinone

whereas the molecular weights of K₁ and K₂ are 450 and 580 respectively Menadione is a yellow, crystalline compound (melting point 106°), slightly soluhle in water and soluble in alcohol, ether, acetone, and glacial acetic acid and in vegetable oils. It is light-sensitive and, in common with other quinoues, has a burning taste and is irritating to mucous membranes. The long substituent side chains of the natural vitamins diminish their solubility in water and, at the same time, their taste.

Many active derivatives and homologs of menadione or the natural vitamins have been prepared but always with a diminution or loss of hiological potency. Phthiccol (2-methyl-3-hydroxy-1,4-naphthoquinone), a constituent of the tuhercle hacillus, was early recognized to have slight vitamin K activity. Substitution of the 2-methyl group by hydrogen or various alkyl radicals results in marked loss of activity. Substitution in the 3 position does not cause a serious drop in potency, although a double hond in the position shown for the phytyl group in vitamin K₁ and a long branched side chain are essential in such substituted compounds.

The hydroquinones of vitamins K_1 and K_2 or of menadione may be converted into diacetates, diphosphates, disulfates, etc., which possess lower activity than the rorresponding quinones but are less irritating to the mucosa of the mouth and gastrointestinal tract. Nevertheless, certain of these derivatives which, unlike natural vitamin K, are water-soluble and more stable to light and air have heen employed chnically. Important among these is menadione sodium bisulfite, a white crystalline powder containing 49 per cent menadione

DETERMINATION OF VITAMIN K

Physical or chemical procedures for the estimation of vitamin K activity are complicated by the presence of interfering substances (including vitamins A and E), the different forms in which vitamin K occurs in nature, and their susceptibility to destruction by light, heat, and air, especially during the saponification and extraction steps. The method of Scudi and Bulls⁴⁷ livelving reduction of the quinones by catalytic hydrogenation, and subsequent reaction of the hydroquinones with 2,6-dichlorophenol indophenol, appears to offer the heat approach to the determination of vitamin K in extracts of natural materials including blood and plasma.

⁴⁴⁷ Soudi and Buhs: J. Biol. Chem , 141, 451 (1941); 143, 665 (1912). 144, 599 (1942).

The extent of the reaction is measured in a photoelectric colorimeter. The use of hydrosulfite has been recommended to avoid destruction of the vitamin during saponification.

ASSAY FOR VITAMIN K

Biological Method of Almquist ** Principle The assay is based on the relative doses of assay material and of standard minadione required to restore the prothrom bin elotting time of the blood of visuam K depleted chicks

Procedure One- or two day old chicks are placed in heated (90° to 95° f.) battery brooders with wire-mesh floors. To prevent consumption of vitamin K from bacterial synthesis, food and water should be provided through apertures outside the cage, and moist or soiled food should be discarded. The following ration is fed ad librum

Sardine meal (ether extracted)	17 5
Dned brewers yeast (ether extracted)	7 ə
Ground polished rice	72 5
Codliver oil	10
Calcium carbonate	0.5
Manganous sulfate	0 000

After 10 to 14 days, when the clotting time of the blood of 5 per cent of the chicks is 15 minutes or more, divide them Into groups of 12 (Determine clotting time as foliows Withdraw a few drops of blood from clean cut of exposed wing sein Piace in small viats and shake in water bath at 38 5° to 39° C Time from withdrawal of blood to formation of a firm clot)

Maintain one group on basal ration as negative controls, at least two groups on different dosages of U S P menadione, and at least one on each product to be assayed Administer doses orally in 0 1 mi water or ethyl laurate, depending on solubility Open mouth of chick by applying pressure at the corners so that the dose may be given with a tuberculin syrings (fitted with a blunt edged needle) well down in the throat Do not permit access to food or water for one half hour alter dosage Repeat dosage for four days at 24 hour intervals

Determine prothrombin clotting time on all chicks 24 hours after ad ministration of last dose, as follows Place 0.2 mi of 0.1 M sodium oxalate in short, narrow tubes graduated at 2 ml introduce 2 ml blood from each chick into a tube (Blood may be obtained by decapitation with scissors and directing flow into tube with fingers) Shake thoroughly Plpte 0.1 ml portions into small flat-bottomed vials (15 × 50 mm) Add 0.2 ml of clotting agent " and start timing with stopwatch Place vials in thermostatically controlled water bath at 38.5" to 39" C so that they are tilted at an angle of 45" in a device to permit moving them to a vertical position once per second When gelatinous film (Colt) covers bottom of vial stop with and vertical position once by the order of the colt of the col

"prothrombin time" Duplicate test on each blood sample until results agree within 2 seconds

Calculation Plot the mean prothrombin time (MPT) against log micrograms of menadione for reference groups and draw best-fitting straight line Interpolate equivalent dosage of assay material (1 μ g menadione = 1 A.O A C unit of vitamin K activity)

THE PHYSIOLOGICAL AVAILABILITY OF THE VITAMINS

The vitamin content of a food, or of the diet as a whole, as determined by chemical or microhiological assay, does not always constitute a rehable indication of the actual amount of utilizable vitamin consumed Failure of foods to be completely digested and absorbed from the intestinal tract, in vito destruction due to oxidation or incompatibilities among the dietary constituents, abnormal pH conditions due to gastric hypoacidity or the use of antacids-these are among the many factors which influence the physiological availability of the vitamins Vitamins often exist in nature in firm union with protein or other compounds, necessitating hydrolysis prior to chemical or microbiological assay to a degree not dupheated in the gastrointestinal tract Animal assays are a hetter measure of physiologically available vitamins, although they do not always simulate conditions in man For example, assay materials sometimes have to he subdivided or dissolved in order to feed the small doses required by animals, moreover, purified hasal rations deviate considerably from the diet of man

These factors emphasize the desirability of a human assay technique for determining physiologically available vitamins in foods and other sources and for determining the effect of various conditions which influence their availability A highly reproducible procedure developed by Meluick, Hochberg, and Oser, 450 is hased on the fact that uormal buman subjects subsisting on a complete diet excrete the water-soluble vitamins in the urine in direct proportion to the quantity consumed above the adequate basal level These vitamins may be excreted either unchanged or as derivatives The hiear relationship of excretion to dosage is established for a group of experimental subjects by feeding the vitamins in pure solution, the form in which they are most completely available Fig 292 illustrates a series of such unuary exerction studies for thiamine, riboflavin, macmamide, and ascorbie acid. To determine the physiologically available vitamin coutent of an unknown material, the experiment is repeated, feeding an amount of the test food furnishing (according to chemical or microbiological analysis) a critical dosc of the vitamin iu question The relation between the extra urinary excretion of the vitamin in the test dose and in the pure solution constitutes the index of physiclogical availability. For example, if under the test conditions threequarters as much extra thiamine (i.e., above the basal excretion level) is found in the 24-hour urme following the ingestion of a given food as was exercted following the ingestion of au equivalent amount of thiamine in the form of the pure solution, the physiological availability of the thiamine in the test food is said to be 75 per ecut

⁴¹⁹ Melnick Hochberg, and Oser J Vutration 30, 67 (1945)

The illustrations of the application of this technique which follow are taken from the work of McImck, Hochberg, and Oser 40 The method may be applied only to those vitamins normally exercted via the urinary tract, which of course excludes the fat soluble vitamins

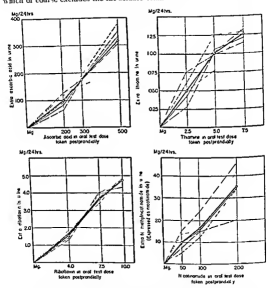


FIG. 292 THE LINEAR RELATIONSHIP BETWEEN DOSAGE WITH THE WATER SOLUBLE VITAMINS ASCORBE ACID THIAMINE RESOFLANT AND NICOTIAM DE AND THE EXTRA UNIVARY EXCRETION OF THE VITAMINS (OR DERIVATIVE).

The test doses in augusus solution (off. 2.0) were taken could unumerisately

The test doses in aqueous solution (pH 3 0) were taken orally immediately after dinner. The fine lines represent the responses of the individual subjects, whereas the heavy line indicates the average responses.

Any suitable method may be employed for the analysis of the urine for the vitamin and its metabolites. It is worth noting that thiamine is excreted in the urine as the free unphosphorylated vitamin, ascorbic acid is largely, but not entirely in the reduced state so that reduction of the dehydro form is necessary intotime and is exerted principally as the metabolite Vi-methylincotinamide, and vitamin Be-theliv as pyridovic acid

14 2

920 ~

230 g

150 g

150 g

12 g

1 orange

2 fried eggs

1 lettuce serving

1 tomato serving

2 shees of bread (enriched)

(bodeward) toward for search of

Determinations of the physiological availability of the vitamins employing this procedure have been reported for pharmaceutical tablets in which the vitamin ingredients are protectively coated to insure their stability, for the thiamine in live yeast cells, for ascorbic acid as influenced by the presence of copper or ascorbic acid oxidase, for thiamine as influenced by the presence of the antithiamine factor in raw fish such as clams, and for the B vitamins in the presence of adsorbing agents

Determination of the Physiolagical Availability of Thiomine in Lie Yeast Cakes Principle. Unless yeast cells are destroyed by autolysis or heat they yield only a small proportion of their thiamine content in the human gastrointestinal tract Subjects in whom nutritional equilibrium has been established each receive, on a control day 6 mg of thiamine in pure solution and in divided doses after each ineal After an intervening period has elapsed the same subjects receive 6 mg of thamine (determined by chemical analysis) in the form of live yeast cakes similarly divided after meals. The ratio of extra thiamine exerction in the test and control days is the measure of physiological availability of thiamine in the verst product.

COMPOSITION OF BASAL DIET

150 g

Breakfast

2 butter squares

1 aloco of mult

I glass of mulk

2 hard sugar candies

1 apple

1 banana

50 g	I glass of milk	200 g
Lun	cheon	
150 g	5 butter squares	35 g
65 g	1 glass of milk	230 g
60 g	1 serving of apple pie	155 g
70 g	2 hard sugar candies	12 g
50 g	_	_
Su	nner	
	150 g 65 g 60 g 70 g 50 g	Luncheon 150 g 5 butter squares 65 g 1 glass of milk 60 g 1 serving of apple pie 70 g 2 hard sugar candies

4 butter squares 28 g

90 c

25 €

70 g

66 g

Analyses Conducted on Aliquots of the Composite Diet				
Proximate Analysis V	alues Found ⁴⁵¹	Vitanun Content	Values Found461	
Total weight	2075 g	Thiamine	1 34 mg	
Total solids	540 g	Thiamine Calorie ratio	0.5	
Moisture	1535 g	Thiamine Nonfat		
Protein	101 g	calorie ratio	0.9	
hat (ether extract)	129 g	Ascorbic acid	115 mg	
\sh	22 g	Riboflavin	2 54 mg	
Crude fiber	6 g	Niacin	22 0 mg	
Carbohydrate (by differen-		i		
Caloric value	2710 Cal	İ		
Nonfat calories	1550 Cal	í		

⁴⁴¹ Expressed in terms of total food consumed in the three meals.

Procedure A group of five subjects in good nutritional status, but not unduly saturated with respect to thiamine, is assigned for four days to a complete dlet (as illustrated in the table above) The dietary schedule is as follows a basal day followed immediately by a control (standard dose) day, then, after an interval of one to two weeks another basal day followed immediately by a test (assay dose) day Just before the largest (noonday) meal of each basal day, the urine is volded and discarded. The urine collection is then begun (in a 2 liter brown glass bottle containing 20 ml 10 per cent sulfuric acid as a preservative) this marking the beginning of the control (or test) period On the control day, 2 mg thiamine, dissolved in water or milk, ls taken after each meal The 24 bour urine is collected as described above At least one week (preferably two) should elapse to provide for complete flushing of this control dose, during which no remaining collections are made and the subjects are free to eat adequate diets of their own selection "The basal diet is then resumed and the urine collected for a 24 hour period. On the succeeding (test) day the same diet is consumed, but in place of the thiamine supplement of the control period each meal is supplemented with a suspension of yeast cake in milk or water furnishing the equivalent of 2 mg of thlamine (It is necessary, of course, to establish the thiamine content of the yeast under test by actual analysis rather than to assume the quantity stated to be present on the label)

Analyses of the 24 hour urine collections made on the basal, control, and test days are performed according to the colorimetric method described on p 1144

CALCULATION From the thiamine excretion of both the control and test days, subtract that of the preceding hasal days if the absolute increment of thinming intake 15 not identical on the control and test days express the increment in thiamine excretion as percentages of the respective supplementary doses. The extra exerction of thismine following yeast dosage divided by that following dosage in the form of pure solution multiplied by 100 gives the percentage physiological availability of the thiamine con tained in the veast

The experiment may be so extended that each test period is allowed to run for 48 instead of 24 hours. In experiments reported by the authors of this method 44 this was done in order to eliminate the possibility of a delayed rate of absorption of thismine from bye yeast cells as compared with pure solutions. That this does not occur 13 illustrated by the data in the table on p 1285

Effect of Dietary Thiaminase in Fish Products 444 Chastek paralysis an acute dietary disease of loxes is caused by including 10 per cent or more of certain species of uncooked fish in the diet and may be cured or prevented by giving adequate amounts of thiamine 411 Thiamine deficiency has been observed in cats led a diet consisting ex clusively of salt-water herring.446 The enzymic nature of the antithiamine factor was suggested by the lact that concentrates possessed protein characteristics and reacted like a typical enzyme toward inhibitor substances "The over-all reaction has been demonstrated to be tile hydrolytic cleavage of the vitamin between the pyrimidine and theazole rings.448

The antithiamine factor has been found in 15 out of 31 species of fresh water fish

⁴³² Pork meats and pharmaceutical vitamin preparations especially should be avoided

in Hochberg Melnick and Oner J Vatration 3a 201 (1945)
in Melnick, Hochberg, and Oner J Vatration 3a 81 (1945)
in Green Carlson and Evans J Vatration 31 165 (1942)

^{***} Smith and Prout Free See Expl Biol Med 56 1 (1944)
*** Smith and Prout Free See Expl Biol Med 56 1 (1944)
*** Sealock and Goodland J Am Chem See 66, 507 (1944)
*** hrampits and Woolley J Biol Chem 152, 9 (1944)

RATE AND DEGREE OF URINADA EXCRETION OF THIAMINE FOLLOWING DOSACE OF THE VITAMIN IN LIVE YEAST CASES AND IN PURE SOLUTION

	Control Period		Test Period			
Subtect	Basal	After 5 76 mg of Thiamine in Solution 459		Basal	4fler 6 Live Yeast Cakes***	
	Excretion	1st 24 hrs	2nd 24 hrs	Excretion	1st 24 hrs	2nd 24 hrs
		Mg per da	,		Mg per day	ı
HH IC EM WH DM	0 20 0 20 0 20 0 20 0 26 0 17	1 56 1 68 1 67 1 78 1 48	0 55 0 59 0 64 0 57 0 45	0 25 0 24 0 28 0 23 0 21	0 44 0 56 0 37 0 70 0 30	0 25 0 37 0 29 0 40 0 22
Average	0 21	1 63	0 56	0 24	0 47	0 31

 $\frac{0.47 - 0.24}{1.63 - 0.21} \times 100 = 16.2$ per cent availability

163 - 021 (as measured by first 24 hrs 'excretion).

 $\frac{(0.47+0.31)-2(0.24)}{(1.63+0.56)-2(0.21)} \times 100 = 170$ per cent availability (as measured by 48 hrs 'excretion)

tested,441 in clams,442 in the Atlantic herring, in whiting, and in the Pacific mackerel, oysters are among the fish not containing this factor. In many parts of the world, fish are eaten raw or only slightly heated (e.g., smoked). The following experiment demonstrates the effect of the thuminase in raw clams in subjects receiving thiamine as a dietary supplement, and the inhibition of this effect by heat inactivation of the destructive enzyme. This experiment may be performed with certain other edible raw or lightly smoked fish.

Procedure. Whip 100 g. raw clams in 400 ml. water in a Waring Blendor. Adjust 100 ml. of this suspension to pH 45, and boil under reflux for 20 minutes. Cool and readjust to original pH. Prepare an aqueous thlamine solution containing 250 pg. per ml. Set up three 100-ml. centrifuge tubes as follows (I) 50 ml. water plus 1 ml. thiamine solution, (2) 50 ml. unheated clam suspension plus 1 ml thiamine solution, (3) 50 ml. heated clam suspension plus 1 ml. thiamine solution.

Store the suspensions for 6 hours at 37° C. with frequent agitation. Centrifuge. To 10 ml. of the clear supernatants, add 10 ml. of phenol-alcohol reagent and complete the colorimetric test for thiamine described on p. 1145.

For the in 1300 availability study, choose five normal subjects with good dietary histories. Feed the basal duet shown on p. 1283, starting at noon, and collect 24-hour urine samples in 2-liter hottles containing 20 ml. 10 per cent

⁴⁴⁹ Taken as three I 92-mg doses of thamme in milk one after each meal

⁴⁴⁶ Containing 5 75 mg of thiamine Two cakes, containing 1 92 mg., suspended in milk and taken after each meal

⁴⁴¹ Deutsch and Hasler Proc Soc Exptl Biol Med , 53, 63 (1943)

Woolley: Personal communication (1943)

sulfuric acid. On the second day, just after the midday meal, feed 5 mg thramine in 30 ml aqueous solution and collect the second 24 hour samples

Before starting the test period allow two weeks to intervene during which the subjects return to their usual, adequate diets. The test period comprises four days. On the first three feed the subjects the basal diet, but at the end of each meal, add a fresh, raw clarn weighing approximately 35 g. (without the shell) which may be swallowed whole. At the beginning of the fourth day (after the midday meal) the subjects ingest 5 mg. thiamine in 30 ml water followed by a clarn. Continue the experiment till the end of the fourth day, feeding a clarn with each meal as before. Collect 24 hour urine samples on all four days.

Analyze the urine samples for thiamline as described on p 1144 Note the prompt and marked decrease in basal urinary excretion values when clams are intested. Note also the effect of the clams on the test dose of the vitamin

VI STUDENT EXERCISES

I Demonstrations of Vitamins $A, B_1 D$ and G Divide at least ax rate 21 to 28 days old into two groups so that there are twice as many rats in one group as in the other Keep the rats in individual cages and follow the instructions for their care outlined on p 1364 ff Feed ad hb one of the following diets to the larger group (experimental) and to the other group (preventive controls) feed the same diet plus one of the indicated supplements daily (except Sundays)

(a) Vitamin A Deficient Diet (see p. 1262) Supplements. One drop U S P codiver oil dally or one μg of β carotene in 0.1 ml. of cottonseed oil, or 3 drops of butterfat

Weigh rate semiweekly and note changes in growth, vaginal smears (usu ally during the fourth week), condition of the eyes, and appearance of the life Compare the experimental group with the control group. Allow half of the experimental group to continue on the basal diet until they die at which time perform autopsies noting the presence of localized infections in the respiratory tract, tongue abscesses renal calcull etc. When growth has definitely cased on the basal diet begin to feed the remainder of the experimental group one drop of codiver oil (or other source of vitamin A). Record your observations.

(b) Thionnine Deficient Diet (see p. 1147). Supplements 0.5 g. dried yeast 30 g. whole wheat flour 5 g.g. of crystalline thionnine diluted in 100 mg. con 6 ectioners' sugar-cornstarch mixture (1 i).

Welfs rate semiweekly Beginning with the third week note signs of polyneuritis Spin the rats by rolling the tail between the hands Convulsive selzures following this treatment are evidence of polyneuritis. When this is observed for two or three consecutive days begin to feed half of the experimental group a thiamine containing supplement. Allow the remainder of the experimental group to continue on the basal diet until death. Plot growth curres and compare with curative and preventive control groups.

(c) Vitamin D Deficiency (see p 1261) Supplements One drop of U S P codliver oil or 0 1 g of irradiated yeast *** or one hour s exposure to sunlight (if the weather is sufficiently warm) or 15 minutes* exposure at a distance

^{**}Irradiated yeast may be prepared by exposing a thin layer ()% inch) of dried brewers yeast at a distance of 18 meles from a quarta in every vapor or carbon are lamp for 20 minutes rating over the surface every 5 minutes.

of 30 inches from a source of ultravioler radiation. Feed the experimental and preventive control groups either of the rachitogenic diets described on p. 1261. Record body weights at semiweekly intervals, and after three weeks ex-

Record body weights at semiweekly intervals, and after three weeks examine each rat for evidence of rickets. Note posture, galt, paralysis of hind legs, enlargement of knee joint, etc. Sacrifice one rat from each group, perform autopsy (note beading of rih cartilage) and line test of the tibia as described on p. 1265. If possible, take roentgenograms of at least one rat in each group (cf. Fig. 289). After definite, gross signs of rickets are observed in the experimental group (between three and four weeks), add a vitamin D supplement to the diet of haif the group for a two-week curative period. Repeat above examinations for rickets. Compare all animals at autopsy. Dissect tibiae free from connective tissue, wrap in marked filter paper, extract for 24 hours with acetone in a Soxhlet extractor, dry in oven, and determine bone ash on "dry fat-free basis."

(d) Riboflovin Deficient Diet (see pp. 1163-4): Supplements. 3 ml. whole milk; or, 0.2 g. drued brewers' yeast; or, 10 µg. crystalline riboflavin.

Weigh rats semiweekly. When growth ceases in the experimental group, continue half of the group on the basal diet and add to the diet of the other balf the same supplement fed the preventive control group. Plot growth curves over a total period of six to eight weeks. Note any signs of alopecia or dermatitis.

2. Demonstration of Vitamin C Deficiency (Scurvy). Divide six guinea pigs weighing about 280 to 300 g, into an experimental group of four, and a preventure control group of two. Feed both groups od lib. one of the scorbutigenic diets described on p. 1239, supplementing the diet of the preventive controls daily with one of the following: 1.5 ml. of fees borange juice; or, a fresh carrot or small potato; or, 0.5 ml. freshly prepared 0.1 per cent ascorbic acid. Liquid supplements should be piptted directly into the mouth; the ascorbic acid solution may be sweetened with cane sugar.

Weigh guinea pigs semiweelly; beginning with the fourteenth day examine daily for signs of scury (see p. 1240) in the experimental group, particularly, sensitive joints, bemorthagic gums, loose teetb, and characteristic posture. These should become pronounced during the fourth or fifth weeks. However, when definite symptoms are noted and growth has stopped, begin to feed half the experimental group the same antiscorbutic supplement which the preventive controls receive. Terminate the experiment after a six weeks' curative period. Autopsy each guinea pig either at death or at termination, comparing the negative controls with the preventive and curative groups. Examine especially for enlarged joints in ribs (beading) and leg bones, subcutaneous and intramuscular hemorrhages, loose, fragile teeth and soft hemorrhagic gums.

- 3. Demonstration of Vitamin E, Place two male (A and B) and two female (C and D) rats at weaning, on the vitamin E deficient diet described on p. 1274 and two males (E and F) and one female (G) on the same diet supplemented with a daily allotment of 3 drops of nonrancid wheat germ oil. At the age of about three months, mate these rats in the following sequence (removing the supplement-fed rats from the cage for their daily dose of wheat germ oil):
 - I. Male E with females C and D.
 - Allow mating to continue until each female shows a typical resorption. (The first impregnation may result in the birth of a litter.) (See IV.)

Stigmas Suggesting Deficiency of Vitamin A

Yerosis of the conjunctiva

Thickening with loss of transparency, so that only the more superficial vessels of the bulbar conjunctiva are clearly seen, associated with more or less yellow pigmentation, especially, along the horizontal mendian of the cyclicall, infrequently associated with small foamlike plaques called Bitot's spot.

Papular eruptions of pilosebaceous follicles

A grater-like feel, which in early stages resembles goosefiesh but, when more fully developed, presents the picture of keratosa plans. The extensor surfaces of the arms and flushs and the flexor surfaces of the legs are primarily affected Verous or auteatons of the sign primarily affected.

Dryness, scaliness and ernking, in extreme cases resembling alligator skin in early stages the condition is associated with keratosis plans; but it persists and extends after folledes have disappeared, the body haus being broken and later lost till parts of the body are involved, but the skin of the extremities, particularly of the legs, is more severely affected than the skin of the head and the trunk

Follicular conjunctivitie

Hypertrophy of the follieles, particularly of the lower cyclids

Ninhi blindness

Conspicuous only in cases of advanced, severe deficiency

heratomalacia

Thickening with subsequent ulceration and necrosis of the cornea present only in most severe and advanced forms of deficiency

Stigmas Suggesting Deficiency of Thiamine

Loss of strength of the quadraceps disproportion ate to loss of general strength, evidenced by difficulty in rising from the squatting position Loss of vibration sense first of the toes and later of the malleol and tibes.

Tenderness of the cultes and hyperesthesia of the

Diminution and loss of the ackilles tendon and patellar reflexes first Other tendon reflexes are lost in the later stages of the polyneuritis of dry beriben

ounta found

Treatment of Vitamin A

Larly deficiency state

25,000 USP units of vita

More chronic states

25,000 USP units of vits min A two to three times daily for a prolonged period

Treatment of Thiamine Deficiency

Acute deficiency state

10 to 20 mg or more of thiamine twice daily until relief of symptoms, this may be days or weeks

Chronic deficiency state

o to 10 mg or more of thusmine twice daily for a prolonged period Enlarged heart with dependent edema and elevated venous pressure. Poor response to rest and administration of digitals unless thaining is given. This is a late manifestation of severe deficiency (beriker) heart).

Papillary edema with refinal hemorrhages associated with ophthalmoplegia and polyneurits. The condition is a late manifestation of severe deficiency.

Stigmas Suggesting Deficiency of Riboflavin

Congestion of the limbic plexus

Visible with a small hand lens or the +20 lens of the ophthalmoscope, invasion of the cornea by capillaries arising from this plexus (vascularization) requires a biomicroscope and sht lamp for detection

Cherlosis

Represented in chronic deficiency by excessive and irregular winkling, in acute deficiency by swelling and erasure of the normal winkling of the lips Reddening, thinning, scaling, chapping

of epithelium are associated

Angular standaties

Various combinations of crythema and open
fissuring in the angles of the mouth with or
without a white, most maceration (perfecbe),
sears of healed fissures

D.ssebacia

An erythema overlaid with somewhat greaxy, flaky accumulations resembing bear frost, noted mostly in the alae nast, canthi, pinnae, and other folds of the skin, accompanied in some cases hy coarsening and elevation of the sebaceous follicles of the nose and cheeks, the latter also seen with deficiency of vitamin A.

Magenta tongue

A purplish red coloring with moderate edema and flattening of filiform papillae, observed in more advanced deficiency

Stigmas Suggesting Deficiency of Niacin

Edema of the tongue

Shown by dental indentations Increased redness of the tonnuc

Beefy red in chronic states, scarlet red in severe

acute deficiency

Congestion and hypertrophy of the papillae of the

tongue, followed by fusion and atrophy
In early stages the fungiform papillae are con
gested and hypertrophied. This is followed by

Treatment of Riboflavin Deficiency

Acute deficiency state
5 mg of rihoflavin three times
daily for weeks

Chronic deficiency state

3 to 5 mg of riboflavin three times daily for a prolonged period

Treatment of Niacin Deficiency

teute deficiency state
100 mg or more of macmain
ide twice daily for weeks

Chronic state

100 mg of macmanide twice daily over a prolonged period 11. Male F with female G.

Allow mating to continue until two live litters are born and carried through the lactation period.

III. Males A and B with female G.

After one month, during which no pregnancy should result, kill male A, weigh the testicles and examine histologically. Compare with normal male E, of the same age.

- IV. Begin feeding male B and female C, 3 drops of wheat germ oil dally at conclusion of previous matings Wate F with C and D. Only C should produce a live litter while D should show resorption.
 - V. Male B with female G.

No live litter should result in spite of the proved fertility of female C. Record your observations and draw conclusions as to the effect of vitamin E deficiency on male and female fertility.

4. Studies on the Planning of Experimentof Diets. With the ald of the data given in the various tables on pp. 1335 to 1363, and of other available reference books from which the calorific value, nitrogen content, vitamin content, etc , may be obtained, plan experimental diets as described below, using the following dietary constituents. Casein, egg albumin, gelatin, gluten, whole wheat, lean beef heart, dried milk, dried liver, starch, sugar, agar, lard, hydrogenated vegetable oil, butter, butterfat, codliver oil, dried yeast, orange julce, tomato juice, sait mixtures (see pp. 1262, 1374, and 1375) etc.

Plan diets which may be predicted to have the following effects, and hand in a report to your instructor explaining the reasons for your selections Include in this report the calorliic value, the protein content, and any other properties of the diet your instructor may suggest

- 1. A dlet that will produce ophthalmia
- 2. A dlet that will promote growth but cause rickets.
- 3. A dlet that will cure or prevent rickets.
- 4. A diet that will prevent growth but not cause rickets.
- 5. A diet that will restore growth following Diet 4
- 6. A diet that will prevent normal reproduction.
- 7 A diet that will cause scurvy
- 8 A diet that will cure or prevent scurvy.
- 9. A diet that will cause ketonurso.
- 10 A diet that will correct this ketonursa.
- 11 A diet that will cause polyneuritis.
- 12 A diet that will cure polyneuritis
- 13. A diet that will be conducive to longevity and the birth and reoring of sturdy offspring
- 14. A dier that will produce o dermotitis in rats
- 15 A diet that will cure this dermatitis 16 A diet that will cause one mua in rots
- 17. A diet that will correct this onemia.

CLINICAL VITAMIN DEFICIENCIES

SYLLABUS OF "STIGMAS, SYMPTOMS AND THERAPY" OF THE COUNCIL ON FOODS AND NUTRITION OF THE AMERICAN MEDICAL ASSOCIATION**

The stigmas and symptoms associated with deficiency of vitamin A, thiamine, riboflavin, niacin, ascorbic acid, vitamin D and vitamin K are listed in this syllabus together with a statement concerning treatment of each deficiency. Deficiencies of several vitamins, notably biotin, pyridoxine, pantothenic acid, and vitamin E, are not accompanied by stigmas which can be recognized at present 455 The subject is in a stage of fluidity and development which probably will necessitate early revision or amplification Particularly is this true of the diagnosis and treatment of deficiency of folic acid. This vitamin has been prepared in isolated form so recently that its consideration here is omitted. Not many of the stigmas listed are diagnostic of a vitainin deficiency in themselves, but the occurrence of several of these stigmas in association is at least presumptive evidence of some nutritional failure. Vitamin deficiencies commonly eucountered in chinical practice are multiple. Scrutiny of the dietary history is indicated in cases in which several of the stigmas listed are present Due attention should be paid to the well known fact that stresses such as pregnancy. exposure or disease may occasion the development of deficiency states when the diet otherwise might be considered adequate.

Treatment for a deficiency involves administration orally or, if need he, parenterally of large enough doses of the vitamin to be of therapentic value and continuation of this treatment for long enough periods to assure a sausfactory therapeutic trial. However, since the diagnosis is necessarily presumptive in many instances, exclusive dependence on specific therapy is justified only infrequently, and base to good treat ment in all cases as a diet planned to be adequate nutritionally and assurance that the diet is caten "" The diet is important fair the education of the patient and as a means of dispensing factors beretofore not isolated which will be contained in the foods of such a diet. Likewise helpful in treatment because of its content of factors not as yet identified as some good source of the vitamin B complex as a whole. Products such as hiewers' yeast or an extract of such yeast, wheat germ, extracts of cereal grasses or of rice bran, crude extract of liver or desiceated liver represent such sources. For a price who cannot take foods or drugs orally ar in whim absorption is poor, crude her extract may be given intramuscularly or even no occasion it may be diluted with sterile isotonic solution of sodium chloride or destrose and administered by vem

44 Handbook of Nutrition A Symposium Prepared Under the Auspices of the Council on Foods and Nutrition of the American Medical Association 1943

444 Jolliffe J Am. Med. Assoc 129, 613 (1945)

⁴⁴ Reproduced through the courtes; of the American Medical Association from J Am Ved Assoc 131, 666 (1946)

hypertrophy of the filterin papillar and later by their flattening. As they atrophy they fuse or mat together with multiple fisating to give a cobhlesione appearance and finally buldness. Vincant's infection of tongue, and fauers, uteration, and pseudomembrane formation may or may not accompany these changes in the more subvanced stages of this deficiency.

Dermalilis

I rythema, rough scaling, with ulceration and formation of bulbe, affecting primarily areas of the skin exposed to light, namely, wrists, ankles, neck, and face, observed only in severe deficiency (pellagra) and then frequently associated with durrine and dementia

I ncephalopathy

Clouding of consciousness, cogwheel rigidity, and grasping, sucking reflexes observed in acute, severe deficiency

Stigmas Suggesting Deficiency of Ascorbic Acid

Redness, edema, tenderness, and Heeding on pres-

Observed in acute or subscute deficiency of moderate severity, sometimes with, but usually without, other signs of ascorbic acul deficiency

Thickening and increased firmness of the gums
With recession and exposure of the base of the
teeth, including recession of interdental papillae
observed in chronic deficiency

Retraction of the gams
Laving protects between gain and tooth
secondary infection and resulting pyorthea observed in chronic deficiency.

Loosening and shedding of the teeth Increased capillary fra plity

Manifested by petechial himitrhages of the skin especially in the fourniquet test observed in more severe acute and subsecut deficiency Lasy brusing spontaneous ecolymous of the skin idopathic hemorrhage into joints and slow healing of wounds observed in severe acute and subscrute deficiency.

Treatment of Ascorbic Acid Deficiency

Acute or subacute deficiency state

100 ing or more of ascorbic acid daily for weeks Chronic deliciency state

100 mg of ascorbic acid three times daily over a prolonged period junctions (beading, rachitic rosary) and bulging forchead (cranial bosses) which respond in treatment are found in infiney and in the rare example of late rickets

Stigmas Suggesting Deficiency of Vitamin K

A tendency to bleeding

Particularly from minor wounds, related to abnormal lengthening of the prothrombin time, developing spontaneously in newborn meants, observed in adults after treatment with discumarol or large doses of salicylates, in advanced disease of the liver with poor excretion of bde, and in discase of the intestine, such as sprue in which vitamin absorption is disturbed

Treatment of Vitamin K Deficiency

In adults

I mg of vitamin K two to three times daily with or without bile (I g of desiccated bile or bile salts)

In neuborn babies

1 mg of synthetic vitamin K intramuscularly daily in oil solution for several days

Deficiency Symptoms

A diagnosis of vitamin deficiency only rarely can be based on symptoms or less significant abnormalities than those which base been listed thus far. However, such symptoms and abnormalities frequently accompany the more specific lesions of deficiency.

Symptoms commonly observed with deficiency of thiamine, also less conspicuously in deficiency of other vitamins, include apathy, letbargy, increased emotional initiability, hypersequitivity to noise and pariful stimul, beadache, yague fears, confusion of thought, uncertainty of memory, asthema, loss of manual dexterity, insomnia, beart consciousness, paresthesia, anorexia, nausea, flatulence, epigastre pain, constipation Photophobia, burning of the eves. Incrimation and evestrain not relieve do ylasses

are encountered in deficiency of riboflavin

Press, 1947

Other abnormalities unrelated to deficiency of any single vitamin but commonly observed in persons who are malnourished are dry, brittle, lack luster, rebellious, so-called staring head bair, a loss of sleekness analogous to the rough coat of malnourished animals, blephantis, spider like telangeteasis of the face, seborher of the face, patchy pigmentation of the face, especially suborbital and circumoral, sinus arrhythmia, bradycardia, tachycardia, low blood pressure, loss of tone of muscles and anemia.

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Metabolic Antagonists and Antibiotics

Few, if any, enzymes or enzyme systems have absolute specificity for a single substrate. For example, the hippuricase of Taka-diastase will not only catalyze the hydrolysis of hippuric acid, hut also that of m- and p-chloro-, bromo-, iodo-, nitro-, and methyl-hippuric acids. The corresponding ortho substituted acids unite with the enzyme, hut hydrolysis of the peptide linkage does not occur and the enzyme is rendered inactive. There are many similar instances in which compounds other than the usual substrate will combine with an enzyme, but do not react further. Such blocking compounds are usually structurally similar to the normal substrate. The enzyme so blocked is unable to combine with its usual substrate, and the reaction catalyzed by the enzyme is slowed or stopped according to the proportiou of total enzyme molecules thus blocked.

Perhaps the simplest example of inhibition related to structural analogy is found in the case of the cuzyme succinic dehydrogenase. This enzyme ratalyzes the oxidation of succinic acid to fumaric acid in the presence of a hydrogen acceptor (see p. 318). The action of succinic dehydrogenase on succinic acid is almost completely inhibited by the presence of malonic acid. The relation hetween these two compounds is

evident from a comparison of their structures:

H₂C·CH₂COOH COOH Succinic acid H₂C COOH COOH Malonic acid

The inhibition by malonie acid is reversible, that is, it may be overcome by the addition of a sufficient amount of succinic acid, in which case the enzyme behaves as though no inhibitor were present. This reversible relation between succinic and malonie acids, together with the obvious fact that these two substances cannot react with each other, implies that both types of molecule are competing for a reactive group on the enzyme molecule itself. This is therefore an example of competitive inhibition. Since the rate of formation of a particular enzyme-substrate complex depends upon the frequency of collisions of substrate molecules with the enzyme surface, which in turn depends upon the concentration of substrate, the extent to which malonic acid will inactivate succinic dehy-

Ellis and Walker: J. Biol. Chem., 142, 291 (1942).
 Quastel and Wooldridge. Biochem. J., 21, 1224 (1927).

drogenase depends upon the relative concentrations of succinic and malonic acids. This relationship defines competitive as opposed to non-competitive inhibition.—

$$E + S \rightleftharpoons ES \rightarrow E + P$$

$$E + I \rightleftharpoons EI$$

where L is the enzyme involved, S is its normal substrate, I is the competitive inhibitor, usually a structural analog, and P is the product. By the mass-action law, the dissociation constant of the enzyme-substrate complex is K.

$$K_* = \frac{[E][S]}{[ES]}$$

and the dissociation constant of the enzyme-inhibitor complex is K_t

$$K_t = \frac{[L][I]}{[EI]}$$

Dividing K_t by K_s , we can get an inhibition constant, K

$$K = \frac{[I][ES]}{[S][EI]}$$

At 50 per cent inhibition

$$[ES] = [EI]$$

and

$$K = \frac{[I]}{[S]}$$

Thus, the effectiveness of a competitive inhibitor depends upon the ratio of concentrations of inhibitor and normal substrate, and not upon their absolute amounts. The relationship of competition applies to a number of well known enzyme inhibitions, including the inhibition of saccharase by fructees, and of xanthine coxidase by adenine.

In considering enzymes which have several substrates (e.g. amine oxidase) it is not always possible to make a sharp distinction between substrates and competitive milibitors. One substrate may be considered to be competing with the other substrates.

Another example of what is believed by many to be an instance of competitive inhibition is found in the action of the sulfonamides on bacteria. In studying the action of sulfamilamide on certain bacteria, Woods' found that the growth inhibiting effect of this compound could be reversed by a substance present in yeast extracts. This substance was believed to be structurally similar to sulfamilamide, so a number of structural analogs were tested for their antisulfamilamide activity. Of these, p-aminobenzoic acid (PABA) proved to be remarkably effective, the structural

^{*} Woods Brst J Exptl Path 21 74 (1940)

tural similarity of this compound to sulfamilamide is evident from a comparison of their structures



On the basis of his results, Woods suggested that p aminobenzoic acid should prove to be an essential growth factor for bacteria, and this was later shown to be the case

Lampen and Jones' related the need of certain organisms for p aminobenzoic acid to its utilization in the formation of folic acid (see p. 1195) Sulfonamides competitively inhibit enzymatic reactions whereby p aminobenzoic acid is built into the folic acid molecule Most organisms. prohably all organisms, require folic acid Organisms that require their folic acid ready made are not hampered by sulfonamides, which interfere with folic acid formation, but growth of such organisms is inhibited by certain structural analogs of fohe acid itself, such as aminopterin Those organisms which require p-aminobenzoic acid for the synthesis of folic acid within the organism are subject to sulfonamide inhibition. which can be released competitively by p aminobenzoic acid or non competitively by folic acid If enough folic acid is present to meet the growth requirement, no inhibition of growth of Streptococcus faecalis will be brought about by sulfonamides Still other organisms exist, including many common pathogens, which are sensitive to sulfonamides, but are not released from growth inhibition by folic acid. In such instances the essential substance synthesized from p-aminobenzoic acid may be a derivative of fohe acid (such as the citrovorum factor p 1201), and fohe acid itself may not be an intermediate in the formation of the essential substance Alternatively, in some organisms p-aminobenzoic acid may have important functions other than folic acid synthesis Each microbial species must be evaluated for its requirement of fohe acid and related substances As an example of a notable difference, Streptococcus faccalis requires folic acid and synthesizes it from p-aminobenzoic acid, whereas Luctobacellus arabinosus breaks down added folic acid and utilizes the components 6

Fluoroacetate is a metabolic analog of acetate Like acetate, fluoroacetate can form an active 2 carbon fragment which proceeds to form a fluorotricarboxylic acid. This product appears to be the actual auti-

Lampen and Jones J Biol Cher: 166 43a (1946)

Aminopterin (4-aminoptero) Iglutamic scid) is the structural analog most antagonistic to folic acid and dimunishes the utilization of formate in purine and pyrimidine synthesis Aminopteria and other folic acid antagonists have been used in the treatment of leukemia. For a full discussion of folic acid antagonists see Petering Physiol Revs 32, 197 (1952)

^{*} hoft Sevag and Steers J Biol Chem 185 9 (1950)

7 Peters Brit Med J Nov 29 1952 1165 Do not confuse fluoroacetate with indoacetate which is not an antimetabolite Iodoacetate is a nonspecific macrivator of sulfly dryl groups the integrity of which is necessary for the action of numerous enzy n

metabolite, which blocks the normal conversion of the acids of the eitric series into α -ketoglutaric acid. The result is accumulation of citrate

There is another type of inhibition of biological systems and its reversal which has not always been recognized as being somewhat different from the type described here. This is the condition in which an inhibition of enzymatic or cellular activity by one substance is reversed-i e, the activity is restored-by the addition of a second substance which is capable of reacting directly with the inhibitor For example, many compounds are known which react with the sulfhydryl group (-SH) Certain of these compounds will mactivate biological systems, and the activity may be regenerated by adding an excess of a second substance which contains sulfhydryl groups, such as eysteine or mercaptoacetic acid. It is to be noted, however, that the substance which overcomes the inhibition is not necessarily a constituent of the biological system being acted upon by the inhibitor, and the reversal of inhibition is essentially one of neutralizing the activity of the inhibitor by reacting directly with it While this type of study may suggest the kind of compounds or active groups which function in biological systems, it is clearly different from metabolic antagonism as described here

In the following pages examples are given of what appears to be metabolic antagonism between structurally related compounds. It must be recognized that in practically all instances the precise mode of action of oven the normal metabolite is relatively unknown, the action of the antagonist, therefore, can only be a subject for speculation. There is, nevertheless, a fundamental similarity in action among all of the substances to be described, in that their inclusion in a normally functioning biological system leads to signs of a deficiency which resembles that produced by a lack of the normal metabolite concerned, and which can be overcome by the presence of sufficient amount of the latter. The substances to be discussed include structural analogs of the naturally occurring amino acids, vitamins, and hormones.

ANALOGS OF AMINO ACIDS

Methionine and Ethionine. In 1938 Dyer's reported that the S-ethyl analog of methionine, called ethionine, appeared to be toxic to rats on a low-cystine, low methionine diet, whereas in the presence of sufficient methionine loss of weight and death did not occur

^{*} Dyer J Biol Chem 124, 519 (1938)

Subsequent work has shown that ethionine blocks several vital functions of methionine (a) transmethylation, (b) incorporation of methionine and other amino acids into protein, (c) formation of cystine, and (d) high potropie action. The antibacterial action of sulfonamides is partially antagonized by methionine, but is augmented by ethionine. The sulfonamides compete with p-aminobenzoue acid in its function as part of a geographic related to the methylation of homocysteme.

The experimental results of Dyer represent probably the first authentic demonstration of metabolic antagonism involving amino acids, in terms of the concept as it has been presented in the preceding pages

Another example of an autimetabolite to methonine is found in the compound known as methonine sulfoximite

Methionine sulfoximine

This substance was first isolated from the hydrolyzate of the protein of wheat flour which had been oxidized (matured) by means of introgen trichloride (Agene). Such flour is tove to dogs producing the disease known as camine hysteria or running fits. The toxic principle was isolated with the aid of chromatographic techniques and was proved to be a derivative of the mino acid methnonine, produced by the action of the introgen trichloride on the methnonine of the wheat-flour protein. The compound has been synthesized and given the name methnonine sulfoximine because of its structure. The compound shows a lugh degree of species specificity, being much more toxic to dogs than to other laboratory animals, but investigations have revealed no evidence of any effect of the treated flour on human beings. However, introgen trichloride is no longer permitted in the United States for maturing flour.

The tone effect of methonine sulformine on dogs can be readily overcome by the addition of methonine to the diet. The growth of certain microorganisms can be inhibited by low concentrations of methonine sulformine, and this effect is reversed by methonine gutamine, and to some extent by methonine sulforde (but not by methonine sulfone of plutamic acid). Thus methonine sulformine appears to be a true antimetribolite but the mechanism of this action is still obscure.

Phenylalanine and Thienylalanine The synthetic compound β 2-thienylalanine is similar in structure to the amino acid phenylalanine excipt that a —CH—CH—group in the benzene ring of phenylalanine is replaced by —S— in thenylalanine

The growth of certain yeasts and bacteria is inhibited by thienylalanine and the inhibition is overcome by phenylalanine. The growth inhibition of L call caused by 100 parts of then lalanme was 50 per cent nulhified by 5 parts of phenylalanine, but tryptophan was almost as effective as phenylalanine Inhibition of yeast growth by thienylalanine was overcome by certain other amino acids in 10 to 30 times the concentration of phenylalanine required. The tubercle hacillus, which is resistant to most growth inhibitors, is susceptible to thienvialanine, as is vaccinia virus, although in the latter case action on the host cell can not be ruled out Studies with \$-3 thienylalanino have indicated that it is a somewhat more effective phenylalanine antagonist than its 8-2 isomer, whereas B-2-furylalanino is less effective Inhibition of weight gain in young or protein-depleted rats can be produced by phenylalanine antagonists, and reversed by phenylalamne but not by tyrosine Halogenated phenylalanines and phenylserine have been shown to be competitive antagonists to phenylalanine in several microbial species. A very important noncompetitive antagonist to phenylalanine is chloramphenicol, one of the broad spectrum antibiotics (see p. 1317)

The enzymes which catalyze the oxidation of phenylalanino to tyrosine, and through further steps to melanin may be blocked by certain analogs. Theinylalanine will block the first step in L. colt, the oxidation of tyrosine by bacterial and mammalian tyrosinases is blocked by N-acetyl. N formyl, or 3 fluorotyrosine. Dimephrine and norepinephrine are mactivated in the hody by several enzymatic mechanisms. One of these, amine oxidase is competitively blocked by ephedrine, amphelamine, and other related compounds. This blockage of epinephrine inactivation is not a complete explanation of the pharmacological actions of these sympathicominents drugs.

Isomeotime and hydrazide is inhibitory to amine oxidase, and 1-isomeotimyl 2 isopropylhydrazine is about 10 times as effective

CH₂
1 Isonicotinyl-2 Isopropyihydrazine

These two compounds, like other basic antitubercular drugs and antibiotics, inhibit the diamine oxidase of mycobacteria. Against mammalian diamine oxidase, the authorotics streptomycin and neomycin have little effect, but the two hydrazides inhibit both the bacterial and the mammalian cuzyme

Other Amino Acids The amino acid L-canavanine, which occurs in jack beans, is a competitive antagonist to arginine and inhibits growth of *E cols*, of certain strains of *Neurospora*, and of lactic acid bacteria Arginase, however, will split urea from canavanine Citrulline and ornithine will release organisms capable of utilizing these substances from canavanine inhibition

One general method of modifying amino acids consists in replacing the carboxyl group by the sulfome acid group, as in the case of sulfamlamide and p-aminobenzoic acid (p. 1290). These aminosulfome acids prove to be growth inhibitory to microorganisms but the inhibition is apparently nonspecific since it may be reversed by aminocarboxylic acids not structurally related to the inhibiting compound Mellwania interprets this as indicating that structural analogy alone is not the sole consideration in

Owen Karlson and Zeller J Bact 62 53 (1951)

** MeIlwain Brit J Fxptl Path 22 148 (1941)

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explaining the action of metabolic antagonists. Indoleacrylic acid, the deaminated α , β unsaturated analog of tryptophan, has heen reported to inhihit the growth of E. coli, and this inhihition is overcome by tryptophan, but apparently not competitively.

The p and L forms of an amino acid would appear to he competitive analogs in those instances where only one of the two forms is metabolized by an organism. While there are numerous instances demonstrating the lack of availability of one of the two enantiomorphs of an amino acid for growth, there are relatively few examples where competitive inhibition has been demonstrated.

There are numerous instances in which naturally occurring amino acids compete with each other; for example, arginine competes with lysine in a mutant strain of Neurospora. This and other examples have been tabulated by Martin. The possibility that competition between naturally occurring substances plays a part in regulating metaholic processes can not be overlooked. Martin postulates, as the "relativity theory" of the hiological world, that natural metabolite antagonisms are fundamental to life by virtue of stabilizing relative concentrations of available nutrilites. In pharmacology and therapeutics, numerous applications are made of the theory of metabolite antagooism. In relatively few instances can competitive relationship be demonstrated; more often, the action is noncompetitive or irreversible.

ANALOGS OF VITAMINS

Thiamine and Pyrithiamine. Pyrithiamine is the pyridine analog of thiamine, obtained by replacing the —S— group in the thiazole ring of thiamine by —CII=CII—; the structural relationship is thus similar to that between thienylalapine and obenylalapine.

Pyrithiamine was found by Robbins¹¹ to be inhibitory to the growth of certain microorganisms which required an exogenous supply of thiamine for growth, and Woolley and White¹¹ showed that the feeding of pyrithiamine to mice produced a severe deficiency which resembled that of thiamine deficiency and which could be overcome by the administration of thamine; one molecule of thiamine was required for approximately 10 molecules of pyrithiamine. This is one of the earliest instances of a deficency disease produced in animals by the administration of a structural analog of a vitamin.

Hollans Proc. Nat. Acad. Sci., 27, 419 (1911)
 Woolley and White J. Biol. Chem., 149, 285 (1943).

Studies on the juhibition of growth of microorganisms by pyrithiamiue have revealed that the substance is inhibitory only in those instances where the organism requires an external supply of thiamine. If the organism is capable of synthesizing thiamine from the constituents of the medium, pyrithiamine is not inhibitory to growth. This emphasizes the point already made that structural analogy per se is not necessarily the sole factor in determining whether competitive inhibitiou will occur. particularly in the complex metabobe processes of living protoplasm. One interesting example has been presented of a strain of yeast which became resistant to the inhibitory effect of pyrithiamine apparently by the development of a system which permitted liberation of the pyrimidine portion of the molecule and its subsequent utilization in the synthesis of thiamine itself.13

Certain other structural analogs of thiamine have likewise been shown to be antagonistic to thiamine; for example, oxythiamine, in which the NH2 of thiamine is replaced by OH. The thiamine molecule is particularly susceptible to such chemical modification, consisting as it does of two reactive portions, the pyrimidine moiety and the thiazole portion.

Pantothenic Acid and Pantoyltaurine. The replacement of the carboxyl group of pantothenic acid by the sulfonic acid group 14 produces a

substance known as pantoyltaurine (or thiopanie acid):

This substance and related compounds, e.g. the amide, have been shown to be antagonistic to pautothenic acid for a variety of microorganisms; growth inhibition caused by pantoyltaurine can be overcome in most instances by pantothenic acid, and the inhibition appears to be truly competitive in nature. As with thiamine and pyrithiamine, pantoyltaurine is not inhibitory to the growth of those microorganisms which can synthesize the pantothenic acid they need.

Pantoyltaurine has uot as yet been shown to produce unequivocal signs of a pantothenic acid deficiency in animals, probably because of the relatively large amounts which would be required to compete successfully with the pantothenie acid present in animal tissues. Some success bas been obtained in the use of pantoyltaurine as a chemotherapeutic agent in rats infected with an organism susceptible to inhibition by pantoyltanrine, but here again the doses required were so large as to indicate little practical application of this fact. The possibility remains open, however, that further studies along these lines may produce more effective substances. As with thiamine, the presence of two ebemically distinct portions in the pantotbenic acid molecule has permitted the synthesis of a

¹⁸ Woolley: Proc. Soc. Expl. Biol. Med., 55, 179 (1944). 14 Snell: J. Biol. Chem , 139, 975 (1944).

variety of structural analogs other than pantoyltaurine, and certain of these appear to be effective in inhibiting the growth of nucroorganisms

in a manner similar to that described for pantoyltaurine

Riboflavin and Analogs Various structural analogs of riboflavin have been synthesized which appear to be metabolic antagonists of the naturally occurring vitamin One of these is isoriboflavin, in which the two methyl groups are in the 5 6 politions instead of in the 6,7 positions as in the naturally occurring vitamin

5 6 Dimethyl 9 (D 1 ribityl) isoallozazine (isoriboflavin)

The onset of the symptoms of riboflavin deficiency can be hastened in rats already on a suboptimal duet by the feeding of isoriboflavin but his substance does not function as a riboflavin antagonist in any bacterial species so far studied Antagonists similarly effective in animal experiments have been prepared by substituting su_ars other than ribose in the 9-po ition Effective both in animals and bacteria is 2,4-diamino-7,8-dimethyl 10-ribityl of 10-diby drophenazine

Large doses of isombofiavin or p-galactofiavin render mice refractory to implants of lympho.arcoma

Pyridoxine and Analogs Deoxypyridoxine produces acrodynia in the mouse and has produced glosaitis cheilosis and seborrhea in human subjects, corrected promptly by pyridoxine. Methoxypyridoxine is an effective antagonist to pyridoxine in chicks but is much less active in rats and is mactive in the mouse, which can form pyridoxine from methoxypyridoxine. A number of other analogs are antagonistic to pyridoxine when tested with yeast

Nicotime Acid and Analogs. Analogs of nicotime acid which appear to act as competitive antigomists of the latter have been obtained by replacing the carboyl group of meetime acid with either the sulfonic acid group or the acetyl group

The analogous amides have also been studied The sulfone acid analog produces inhibition of growth of certain microorganisms which can be overcome by meetinic acid, but is apparently without effect on animals As with certain other inhibitors already discussed, pyridine sulfonic acid is without effect on those species of microorganisms which can synthesize nucetinic acid again suggesting that cell penetration or local concentration may be as important as simple structural analogy in explaining the action of metabolic antagonists. The acetylpyridine analog of meetine acid brings about reversible signs of meetine acid deficiency in mice and dogs, but is without effect on those microorganisms which have been studied.

Ascorbic Acid and Guccascorbic Acid It was shown by Woolley and Krampitz¹³ in 1943 that a synthetic compound called glucoascorbic acid produced scurvylike symptoms when fed to mice on a highly purified diet. These symptoms could not be cured when ascorbic acid was also included in the diet but disappeared when the glucoascorbic acid was omitted. It will be recalled that mice do not require ascorbic acid in the diet, apparently being capable of synthesizing the amount they niced In the case of the guinea pig however, which develops scurvy on diets free from ascorbic acid. Woolley has reported that a disease which is not exactly like scurvy but which has some similarities to it is produced on diets containing glucoascorbic acid.

¹⁸ Woolley and Krampitz J Expd Wed 78 333 (1943)

effect. The structural relationship between ascorbic acid and glucoascorbic acid is cyident from the following

It appears likely that the relation between ascorbic acid and glucoascorbic acid is one of metabolic antagonism, although the evidence is more limited than for most other examples of this phenomenon.

Vitamin K and Analogs. A himorrhagic disease occurs in cattle from cating spoiled sweet-clover hay The substance responsible is discounsard (see formula), which is an antagonist of the K vitamins, interfering with their utilization in the synthesis of prothrombin. Discounsard and soveral related compounds have been used extensively for the prevention of thrombous. Derivatives of 1,3-indandone have been investigated for anticoagulant activity, and successful clinical use has been reported for the pheny and the diplicative directions.

Bliotin and Analogs. The structural relation to biotin of some analog or biotin which have been shown by Dittiner and du Vigheaudit to have antibiotin activity is as follows:

Blotin sulfone

Evidence for competition between hiotin and its structural analogs has been obtained largely by the use of yeast and bacteria. Inhibition of growth of microorganisms by the three structural analogs whose formulas are given is overcome in each case by the presence of an excess of hiotin, but the three differ among themselves in antihiotin potency and likewise show some differences with different microorganisms. Dethiobiotin for example inhibits the growth of L. casei but for yeast is a growth stimulant, i.e., it replaces biotin. It is probable that this is due to the ability of the yeast cell to convert dethiobiotin into biotin. Biotin sulfone is inhihitory to the growth of both yeast and L. casei, as is imidazolidone caproic acid, and this inhibition is reversed by biotin. It is interesting to note that all of the antibiotins mentioned combine with avidin.

Another group of synthetic structural analogs of biotin has been described by English, et al 17 The most potent in the group, in terms of the inhibition of growth of L. casei, has the following structure:

γ-3,4-Ureylene-cyclohexyl-butyric acid

¹⁷ English, Clapp, Cole, Halverstadt, Lampen, and Roblin: J. Am. Chem. Soc., 67, 295 (1945).

Other members of the series differ from the structure shown in the type of main ring system and the length of the side chain Practically all of the compounds described act as antihotins against either yeast or L cases, but with varying effectiveness, and in every case the inhibition of growth is overcome by the presence of sufficient hiotin

ANALOGS OF HORMONES

Relatively little is known concerning the existence of structural an tagonism among the various hormones of known structure. The various protein hormones are of course out of consideration in this connection because their structures are not known, the concept of an "antihormone" for a protein hormone has an immunological connotation rather than one of structural analogy. Among the hormones of known structure, it has been pointed out by Woolley that the naturally occurring estrogens and androgens are structural analogs of each other, and it is suggested that the known instances of biological antagonism between these two groups of compounds may be related to structural similarity and competitive properties. Woolley has also cited as an instance of structural antagonism the interesting findings of Kuhni's in connection with the sex hormones of algae. Kuhn showed that the two sex hormones were not different compounds but cas and trans modifications of the same compound. The factor determining sex was the ratio between these two forms.

It appears likely, therefore, that as further knowledge is gained in the field of hormone chemistry and physiology the existence of competitive inhibition and modification of biological action by structural difference may become as firmly established for some of the hormones as it now i for vitamins and specific amino acids

ANALOGS OF OTHER METABOLITES

In addition to the specific examples which have been cited of metabolic antagonism between naturally occurring metabolites and synthetic structural analogs the concept has been applied to other classes of compounds of biological importance such as purines porphyrins choline and related compounds amines etc. A detailed consideration of these appears un profitable at this time, references to curtain of them will be found in the reviews by Welch and by Woolley cited in the Bibliography and in Martin's monograph

Successful antidotes for poisonous substances must be able to compete with the receptors in the body which are damaged by the poison. Examples are the counteraction of arsenic poisoning by 23-dimercaptopropanol (B.V.) and the prevention or reversal of eyamide poisoning by vita min Bi_{1...} This substance (hydroxocobalamin) reacts irreversibly in the animal body to form vitamin Bi₁ (syanocobalamin).

ANTIBIOTICS

The terms antibiosis and antibiotic powers were coined by Paul Vullemin in 1883 and vere used by him in a broad sense, referring to all in-

huln ingew Chem \$4,1 (1910)

^{*} Mushett helley Bexer and Liekarls I or Sor Exitl Biol Med 21 234 (1952)

stances where one species destroys the life of another to preserve or maintain its own. One example of antibiosis cited by Vuillemin fits exactly, however, the current and more restricted concept. He mentioned the observation that products secreted by the blue pus microbe would hold the anthrax bacillus in check. He went on to predict that by an understanding of symbiotic and antibiotic powers man would learn to dominate disease. Present-day definitions characterize antibiotics as orgain chemical substances which are produced by microorganisms and have the capacity in dilute solutions to inhibit the growth of other microorganisms and in some situations to destroy them. Opinion is divided whether or not to classify as antibiotics antibacterial substances of animal or vegetable origin.

The blue pus microbe, Pseudomonas aeruginosa, the first microorganism in which the property of antibiosis was clerily recognized, actually produces several definitely identified ocompounds with antibacterial activity. The crude mixture hid been tried half a century ago against anthrax infections and had been found too toxic to the host to be of any value in treatment. It was not until 1939, when Dubos reported on tyrothricin (see p. 1314), that interest in antibiotics was again accentuated.

Unlike the common disinfectants and antiseptics, which usually act by irreversibly denaturing or precipitating protein, or by inactivating functional groups (sulflydryl, iron, etc.) within the cell, the action of antibotics must be much more subtle since they are apparently harmless in the organisms which produce them but inhibitory or tone in the susceptible organism. Most autibioties are primarily bacteriostatic in their action, that is they do not appear to inhibit the metabolic processes of treated microorganisms except where subdivision and growth are concerned. In some instances, however, substrate utilization may be affected. Some antibiotics are bactericial, irreversibly destroying the metabolic processes of the susceptible cell. In some cases concentration may determine whether the action is bacteriostatic or bactericial.

Antibiotics differ considerably in their relative effectiveness against various species of microorganisms, and even against different strains of the same species. In some instances susceptibility to an antibiotic may be considerably modified by variations in the conditions of culture.

The differential susceptibility of various species of microorganisms to a given antibiotic permits the construction of a bacterial "spectrum" for the autibotic, in which the relative effectiveness is determined for a standardized series of test organisms. Such a spectrum may be useful in comparing two supposedly identical or dissimilar autibiotics and in comparing synthetic compounds or derivatives with the naturally occurring autibiotic.

Antibiotics also differ considerably in their relative to airmals and in their effectiveness in vito as compared with in vito. Effectiveness in vito may be modified by the presence in the host of means for metabolizing or destroying the antibiotic

²⁰ Wells J Biol Chem 196 331 (1952)

The Penicillins. In 1929 Fleming reported that a certain species of mold, later recognized to be a strain of Penicillium nodatum and first encountered as an accidental contaminant of hacteriological plate cultures, produced a soluble diffusible substance which inhibited the growth of the common Staphylococcus aureus and certain (hut not all) other microorganisms in culture media. Fleming was able to concentrate this substance from cultures of the mold, and called the material penicillin. It is now known that the original penicillin was a mixture of several similar substances which are called as a class the penicillins.

Fleming showed that his penicillin preparation could be used for the differential separation of resistant and nonresistant species of microorganisms in culture media, and that it was nontoxic to animals. He suggested its possible use in the treatment of infections due to penicillin

sensitive organisms.

These observations however attracted little further attention untiabout ten years later, when Dubos¹² published his striking results on the antibacterial properties of tyrothricin (see p. 1314). It is probable that these results contributed significantly to a reawakening of interest in the antibacterial possibilities of penicillin. In 1940 Chain, Florey, et al., reported the preparation of penicillin in impure form, with some studie on its toxicity to animals and therapeutic value in experimental infections in mice. This was followed by a more extensive study on method for large-scale production, partial purification, and assay, and furthe studies on toxicity and therapeutic value in animals and man. Similas studies on the clemotherapeutic value of penicillin were reported be Dawson, Hobby, Meyer, and Chaffee¹⁴ at ahout the same time. Develorments since then have made penicillin by far the most widely used an important of all the antibiotics thus far known.

Of the various known penicillins, the most important at the presentime is the one named penicillin-G (penicillin-H in Britain). The structul

of penicillin-G is as follows:

names which include the name of the major differentiating group (the R group in the table below) According to this suggestion, penicillin G becomes benzylpenicillin, penicillin-X is p-hydroxybenzylpenicillin, etc

As a class the penicillus are moderately strong monobasic organic acids, soluble in water and in organic solvents such as alcohol, ether, amyl acetate, etc The sodium salt, which is quite soluble in water and alcohol, is the usual commercial form. Neutral aqueous solutions of the sodium salt will retain their activity for several days if stored in the cold. acid, alkaline, and alcoholic solutions rapidly become mactive

THE NATIONAL PENICIPLINS.

	7 ype Structure R-CO-\H-C	H,0S\-C00H
A ame	R	Vame of R
Penicilin F Dihydropenicilin F Penicilin G Penicilin K Penicilin X	CH, CH CH=CH CH;— CH; CH; CH; CH; CH;— CH; CH; CH; CH; CH; CH; CH;— CH; CH; CH; CH; CH; CH;—	Δ² pentenyl n amyl benzyl n heptyl p hydroxybenzyl

All of the penicilin available commercially is obtained from natural sources, i.e., from cultures of mold, of which special strains of P notatum and P chrysogenum have been most commonly employed About 30 other variants of penicillii have been produced biosynthetically by the growth of Penicillium in the presence of specific organic acid or amide precursors 25 These penicilius are all aotibacterial in vitro, and some of them have been used therapeutically, with particular value in patients who have developed sensitivity to benzylpenicillin

Penicillus inhibit and in higher concentrations kill actively growing cells of susceptible strains. Inhibitory or lethal action on resting or non multiplying cells is insignificant By the use of radioactive penicillin, fixation of penicillin by a susceptible organism (Micrococcus puogenes var aureus) was demonstrated to be chiefly in the bacterial cell wall 26 Penicillin does not enter yeast cells, which are unaffected by its antibiotic activity Staphylococci can be grown in high concentrations of penicillin, provided these concentrations are low in proportion to hacterial mass Under these cultural conditions, the conversion of amino acids to bacterial protein was inhibited 27 Instead, extracellular polypeptides were produced in approximately equivalent amounts Peucillins also block the uptake of glutamic acid by susceptible staphylococcal strains during active growth. The formation of a precursor of pentose nucleic acid is also delayed. None of these effects has been clearly established as an example of antagomsm to a specific metabolite. The structural similarity

²¹ Behret's Corse Edwards Garrison Jones Soper Abeele and Whitel ead J Biol Chem 175 703 (1948) Volum biliacs and Felsenfield J Am Med Assoc 143 "94 (1950) ²¹ Few Cooper and Rowles Val re 169 283 (1952) ²² Hotelihas J Exp Med 91, 351 (1950)

of benzylpemeillin to glutathione has been noted, 2 but the evidence that penicillin may act as a clutathione antagonist is scant and inconsistent

Penicilin may be administered orally or by injection, usually intra muscularly Questions of dosage, form of administration, and choice of penicillin as compared with other therapeutic agents will not be con sidered here The monograph of Welch and Lewis gives the hacterial "spectra" and the details of therapeutic application of penicillin and other antibioties Pemeillins are nontoxic to animal tissues in any except very large doses, for example, in ordinary concentrations they have no demonstrable effect on the metabolic characteristics of those isolated animal tissues which have been thus far investigated. They differ among themselves however, in their action on microorganisms, penicilhn G, for example is most effective clinically, while penicillin-K is much less useful Lack of recognition of this fact led to some confusion concerning the therapeutic effectiveness of commercial penicillin preparations during the early stages of their clinical application A wide variety of organisms mostly Gram positive but including some Gram negative, are susceptible to the action of nenicillin

Gramicidin and Tyrocidine The antibiotic preparation obtained by Duhos from cultures of B brevs and called tyrothricin proved on further study to consist of two separate compounds, called gramicidin and tyrocidine They may be separated from tyrothricin by treatment with acctone-ether mixture, in which gramicidin is soluble but tyrocidine is insoluble Gramicidin may be recrystallized from acctone, and tyrocidine by drochloride from acidified alcohol Both of these substances are polypeptides of fairly low molecular weight but as yet unknown structure. They are resistant to the action of proteolytic enzymes. On hydrolysthey yield mixtures of amino acids, all of which have presumably bendentified (see below). It is interesting to note that certain of these amino-acids are of the D configuration, in contrast with the hydrolytic products of animal proteins where the L configuration prevails.

Grammedan is insoluble in water, slightly soluble in ether, and readily soluble in acetone and alcohol Estimates of its molecular weight vary on the basis of the amino acid composition a value of approximately 2800 has been proposed On hydrolysis grammedin yields five known amino acids and the hasic compound ethanolamine These components of the grammedan molecule and their estimated molecular ratios, are as follows n-leuene, 6 L-tryptophan 6 nL-valine 5, L-alanine 3, glycine 2 ethanolamine, 2 making a minimum of 24 residues present The valint was probably racemized during the hydrolysis if it were all present if the property of the property of the property of the property of the property of the property of the property of the property of the enzyme n-deaminase (n-amino acid sound present in grammedin by the use of the enzyme n-deaminase (n-amino acid sound present in grammedin to the property of the property of the property of the property of the property of the carboxylor amino groups so that a cyclic structure is probable.

Gramiedin is inhibitory to all Gram positive organisms except acid fast bacilli but is entirely inactive against Gram negative bacilli. It doe not destroy the respiration of susceptible cells in particular instances the

[&]quot; Grevenstuk Science 114 74 (1941)

[&]quot;Lipmann Dubos and Hotchkiss J Biol Chem 141 163 (1941)

may be highly hacteriostatic but not at all hactericidal, thus indicating that it acts rather as a metabolic inhibitor than as a protoplasmic poison. The therapeutic value of grammadin is limited by its relative toxicity to animals and its low solubility in water, penicillin for example being much superior in both respects. It has, however, found some application in the tonical treatment of infected wounds.

Tyrocidue is a polypeptide containing free amino groups (probably the \(\bar{\partial} \)-amino group of the oriuthine present, see below) and forms a crystalline hydrochloride, which is insoluble in water, acctone, and ether but soluble in alcohol, from which it may be crystallized. On hydrolysis it yields the amino acids phenylalamine, leucine, proline, valine, tyrosine, orintlune, glutamic acid, aspartic acid, and tryptophan Of these, the phenylalamine has the n configuration while all the rest (with the possible exception of tryptophan) have the common is configuration. It is interesting to note that the amino acid valine occurs in one configuration in tyrocidine and in the opposite configuration in gramicidin. The proof of the existence of orinthine in the tyrocidine molecule is probably the first instance of the discovery of this amino acid as a primary constituent of polypeptide chains in nature.

Tyrocidine is strongly bacteriedal for various Gram positive and some Gram negative organisms in title It is mactivated in vivo by blood plasma and hody fluids in general, and so contributes httle to the clinical value of tyrothricin. The importance of tyrothricin in medicine is indicated by its juclusion in the United States Pharmacopeia XIV. Its clinical applicatious are limited, however, by its toxicity, which prevents systemic use. The chief method of administration is by local application.

or by justillation into hody cavities

Streptomycin. This antihotic, obtained from cultures of the actino mycete Streptomyces griscus, was first characterized by Schatz, Bugie, and Waksman in 1944. Its discovery was the result of a systematic study of organisms antagonistic to Gram negative bacteria.

Streptomyon is a water-soluble, erystallizable, organic base, with the structure shown in the formula on page 1316. Its solutions are administered by injection, since absorption from the gastrointestinal tract is minimal.

Although streptomyem was originally introduced as a therapeutic agent against infection with Gram-negative organisms in general, the so-called broad-spectrum antibiotics (which will be considered later) have replaced many of its earlier uses. Against the tubercle bacillus, however, streptomyem and a derivative, dihydrostreptomyem, are toxic, affecting the eighth cranial nerve. Streptomyem tends to cause damage to vestibular function and dihydrostreptomyem to auditory function. Use of a mixture of the two has minimized damage.

Concerning the effect of streptomy can on bacterial metabolism, observations with E coll show that the antibiotic inhibits a condensation between oxalacetic and pyruvic acids which piecedes their oxidation by a path differing from the usual eitre acid cycle Certain variants of E coll which have been made resistant to (or dependent upon) streptomycin

do not utilize these metabolites appreciably, but depend upon other metabolic reactions ¹⁰ Numerous other possible mechanisms of ant bacterial action have been noted

Neomycin, from Streptomyces fradiae, is a mixture of water-soluble basic substances, effective against the tubercle hacillus along with a wide variety of Gram positive and Gram-negative organisms. Its elinical use is limited by a toxic action leading to focal necrosis in the renal tubules.

Bacitracin, from Bacillus lichemformis (closely related to B subtilis) is a polypeptide, comparable in a general way with certain other polypeptides of microbial origin—polymyxin, subtilin, and eumyein—which also are antibiotics Bacitracin has had wide application in the treatment of skin infections, one of its advantages being that it seldom induces allergic reactions. All of the antibiotic polypeptides show some degree of toxicity to the kidney, which limits their systemic use

Broad-spectrum antibiotics. This group of antibiotics is characterized by activity against a great number of different pathogene microorganisms, including certain nekettisiae and large viruses Chloreteracycline (Aureomyen) and oxyletracycline (Terramyen), from Streptomyces aureofaciens and rimosus, respectively, are quite similar in structure Both are relatively nontoxic and are commonly given by mouth although intravenous use is possible.

²⁶ Umbreit Smith and Oginsky J Bact 61 595 (1901) Umbreit Trans N Y Acad-Sci Ser II 18, 12 (1952)

Chloramphenicol was originally identified as a naturally occurring antibiotic, elaborated by a species of Streptomyces. It is now produced synthetically. Chloramphenicol is the first nitrobenzene compound to be found in living cells.

It is a noncompetitive antagonist of phenylalanine. Woolley²¹ has emphasized that noncompetitive antimetabolites are more effective antibiotics, since their activity is not canceled by a simple increase in concentration of the specific metaholite. Chloramphenicol may be given orally or intravenously, and is of particular therapeutic value in typhoid fever and against certain rickettisiae, spirochetes, and viruses. Some degree of calculated risk is involved in its use, since depression of bone marrow function has been reported in a small fraction of cases treated with this agent.

¹¹ Woolley: J. Biol. Chem , 185, 293 (1950).

All three of the broad-spectrum antihotics have the property in common of mhibiting the synthesis of protein in susceptible bacterial cells Chlortetracycline will dissociate nhosphorylation from oxidation in animal tissue or mitochondrial preparations, and at higher concentration will inhihit both processes. At similar concentrations, terramycin shows these actions only minimally 12

Antibiotics in Feeds. The crude residues of Streptomyces aureofaciens and of other organisms grown for the production of antibiotics were introduced into the rations of poultry and pigs as a source of vitamin B12 The growth promoting effect of these antibiotic residues was found to be greater than could be accounted for by their B12 content A similar impetus to growth of pigs, chicks, and poults could be observed on the addition of peniciliin, streptomycin, chlortetracycline. or oxytetracycline to a diet adequate by all recognized standards. The hypothesis that these antibiotics are themselves essential growth factors is unlikely on account of the diversity of their chemical structure, and because a similar effect can be observed upon administration of sulfonamides, phenylarsonic acid and other nonbiological antibacterial agents. It is now understood that the growth stimulus depends upon the more or less selective antibacterial action of these substances, which permits the multiplication of certain bacteria which produce B vitamins and possibly other growth factors, meanwhile suppressing other organisms which would compete with these beneficial organisms or even destroy the growth factors produced by them In some instances actual infections, previously unrecognized, have been shown to be suppressed by the antibiotics included in the rations

DETERMINATION OF PENICILLIN

1. Method of Vincent and Vincent " Principle A disk of filter paper is saiurated with the solution to be analyzed which has been diluted if necessary so as to contain approximately I unit per ml The disk is placed on a outrient agar culture plate impregnated with the test organisms and incubated at 37° C If the solution contains penicillin after incubation a clear zone of inhibition of colony formation surrounds the disk The diameter of this zone is measured and the penicillio content determined by reference to a calibration curve showing the relation between iohibition zone diameters and known amounts of penicillin

Procedure The test organism (Staphylococcus aureus II is recommended) is transferred from an agar slant twice through peptone broth for 24-hour periods at 37° C The second transfer Is held at 5° for 16 to 18 hours

Pipet 10 ml of nutrient agar into uniform flat-hottomed petri dishes and incubate at 37° C. for 16 to 18 hours Transfer to a refrigerator for at least one hour. When the plates are ready, flood each one with 1 ml. of the refrigerated culture of test organisms Remove the excess with a capillary plpet Place the plates at 37° C for one hour to dry At this point it is necessary to use wooden racks which support the top half of the petri dish above the hottom half so that there is about 1/2 inch clearance When the plates are dry, store in an Inverted position in the refrigerator for at least one hour.

²¹ Regns Trans A Y Acad Sci Ser II t5 12 (1952)

²² Regns Trans A Y Acad Sci Ser II t5 12 (1952)

²³ Regns Trans A Y Acad Sci Ser II t5 12 (1944) Similar methods

were described dimont amultaneously by Epstem Folloy Perrins and Lee July 1944) Med 29, 319 (1944) and by Sherwood Falco and de Beer Science 99, 247 (1944)

Sterilize the filter-paper disks³⁴ by dry heat and immerse them for 30 seconds in the solution to be tested. Remove each disk from the fluid with sterile forceps, shake off excess fluid, and place flat on the seeded agar plate. Three disks, evenly spaced, may be placed on a plate, and it is suggested that the plates be run in triplicate for greater accuracy. One of the disks on a plate may sure as a standard peniciliin control.

Incubate the prepared plates (not inverted) at 37° C. for 14 hours. The plates should be placed on a wooden block in the incubator to prevent excess condensation. After the incubation

period, measure to the nearest mm.
the diameter of the zone of inhibition around each disk. Mayerage the

results for each unknown

CALCULATION The penicillin content of the sample is equal to that of a standard solution of penicillin which gives the same diameter of inhibition zone under comparable conditions. This is best established by reference to a previously prepared calibration curve relating known amounts of penicillin to zone drameters To prepare such o curve, set up a series of solutions of penicillin in amounts ranging from 4 to 0 1 units per rul, and carry out the procedure as described for the unknown on each of these solutions Plot a curve with zone diameters in mm on the y axis against units of penicillin per ml on the z axis A curve similar to that

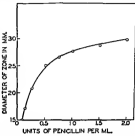


FIG 293 TYPE OF CALIBRATION CURVE OBTAINED IN PENICULIN DETER-MINATION BY METHOD OF VINCENT AND VINCENT

shown in Fig 293 should be obtained It is advisable to check this curve at intervals with standard penicillin solutions, and to reconstruct it if necessary

Interpretation. This method is a slight modification of the original method of Chain, Florey, et al (loc cit) developed at Oxford in England and called the "Oxford cup" method, in which the solution to be tested is placed in small glass cylinders partially immersed in the seeded agar, this solution then diffuses out beneath the cup rim into the agar to produce a zone of inhibition similar to that described here A unit of penicillin (formerly called the Oxford unit and now known as the Florey unit) was defined as that amount which produced a zone of inhibition 24 mm across under the conditions of the cup assay Such a unit is clearly subject to variation from laboratory to laboratory and even in the same laboratory unless assay procedures are rigorously controlled. With the availability of pure crystalline sodium penicillin-G, it is possible to define a penicilin unit in terms of this substance. The Federal Food and Drug Administration of the United States has adopted the following relationship 1 mg of pure sodium penicillin-G is equivalent to 1667 umts of penculin Thus I unit equals 0 6 µg of the sodium salt

The disk method described here has been found to give satisfactory

^{**} Schleicher and Schull No 470 1/2 mch in diameter

³⁴ A bacteriological colony counter equipped with a magnifying glass and a built-in millimeter scale is very satisfactory for this measurement

results on blood serum, spinal fluid, urine, and other solutions whose penicillin activity is to be established. For greatest accuracy the solution to be tested should contain between 0.1 and 4 units per ml

2. Other Methods. Rammelkamp¹⁸ has discribed a procedure for penicilin assay in which the fluid to be analyzed is incubated at various dilutions with a standard inoculum of hemolytic streptococci in the presence of crythrocytes Dilutions of a standard penicillin solution are treated in the same way From the relative dilutions of sample and standard required to inhibit the hemolytic action of the test organisms, the penicillin content is obtained This method, although not so accurate as the one described above, has given satisfactory results on whole blood, crythrocytes, urine, spinal fluid, exudates, and joint fluid Hiscoxii has described a colorimetria procedure, applicable to benzylpenicillin only, for the assay of batches of commercial penicillin This method involves mitration of the benzene ring, reduction, diazotization, and coupling with a color reagent A turbidimetric method has been reported which is applicable to the measurement of any of the commonly used autibiotics in blood or body fluids

For techniques of streptomycin assay, and of tests of organisms for sensitivity to antibiotics, consult the Diagnostic Procedures and Reagents of the American Public Health Association Sensitivity tablets and disks impregnated with varying concentrations of different antibiotics are available from biological supply houses, and each manufacturer recommends procedures and media for use with his product

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Appendix

I. REAGENTS AND SOLUTIONS

REAGENTS FOR PARTICULAR METHODS ARE DESCRIBED IN THE TEXT AND FOOTNOTES FOR THESE METHODS.

SEE INDEX

Acid Digestion Mixture. To 50 ml. 5 per cent copper sulfate solution add 300 ml. 85 per cent phosphoric acid and mix. Add 100 ml. concentrated sulfuric acid (NH₂-free) and mix. Keep in a glass-stoppered bottle, protected against the absorption of ammonia from laboratory atmosphere

Alcohol-Ether Mixture. Mix three volumes 95 per cent redistilled

alcohol and 1 volume redistilled ether.

Alizarin. A 1 per cent solution of alizarin monosodium sulfonate in water.

Alkaline Pyrogallate Reagent. Prepare a solution of potassium hydroxide by dissolving 100 g. in 130 ml. water. In 200 ml. of this solution dissolve 10 0 g. pyrogallic acid.

Almén's Reagent. Prepare by dissolving 5 g. tannic acid in 240 ml.

50 per cent alcohol and adding 10 ml. 25 per cent acetic acid.

Aluminum Hydroxide Cream. To a 1 per cent solution of ammonium alum at room temperature add a slight excess of a 1 per cent solution of ammonium hydroxide. Wash by decantation until the wash water shows only the faintest trace of residue on evaporation. For the preparation of other forms of alumina see p. 329.

o-Aminobenzaldehyde Reagent. Mix 3 g. crystalline o-nitrobenzaldehyde with 50 g. crystalline ferrous sulfate. Add 75 ml concentrated ammonia. Heat on a steam bath for one hour. Distil off the o-aminobenzaldehyde with steam. The mixture before distillation will

keep for two weeks.

Ammoniacal Silver Solution. Dissolve 26 g. silver nitrate in about 500 ml. water, add enough ammonium hydroxide to redissolve the precipitate which forms upon the first addition of the ammonium hydroxide,

and make the volume of the mixture up to 1 liter with water.

Ammonium Molyhdate Solution See Molybdate Solution, p. 1328
Ammonium Thlocyanate Solution. This solution is made of such a strength that 1 ml. of it is equal to 1 ml. of the standard silver nitrate solution mentioned below. To prepare the solution dissolve 12.9 g, ammonium thiocyanate, NH₃SCN, in a little less than a liter of water. In a small flask place 20 ml. of the standard silver nitrate solution, 5 ml. of a cold saturated solution of ferric alum, and 4 ml. nitric acid (sp. 1.2), add water to make the total volume 100 ml., and thoroughly mix the contents of the flask. Now run in the ammonium thiocyanate solution from a buret until a permanent red-brown tinge is produced. This is the end reaction and indicates that the last trace of silver nitrate has been precipitated. Take the huret reading and calculate the amount

1321

of water necessary to use in diluting the ammonium thiocyanate in order that 10 ml of this solution may be exactly caual to 10 ml of the silver mitrate solution Make the dilution and titrate again to be certain that the solution is of the proper strength

Antifoaming Oil Mixture (Use like caprolic alcohol) To 1 volume

erude fuel oil add about 10 volumes tolucne

Asbestos for Suction Filters The asbestos is shredded placed in a wide-mouth flask, and covered with 10 per cent HCl Heat on a water bath for five hours Filter on a Buchner funnel, wash free from acid, return to the flask, cover with 3 per cent \aOH, and heat on a water bath for three hours Tilter, wash free from alkali, then with dilute acid and finally with water until free from acid Suspend in a large volume of water, allowing it to settle for five minutes Pour off the upper two-thirds and discard Repeat the washing of the desired coarse portion several times until the supernatant bound remains nearly elear

Barfoed's Solution. Dissolve 133 g neutral ervstallized copper acetate in 200 ml water, filter if necessary, and add 1 8 ml glacial acetic

hise

Barfoed's Solution (Tauber-Kleiner Modification). Dissolve 24 g copper acetate (Merck, normal, e p) in 450 ml boiling water If a precipi tate forms, do not filter Immediately add 25 ml 8 5 per cent lactic acid (Mallinckrodt, USP, 85 per cent) to the hot solution Shake, nearly all the precipitate will dissolve Cool, dilute to 500 ml . and after sedimenta tion filter off the impunities

Baryta Mixture. A mixture consisting of 1 volume of a saturated solution of banum nitrate and 2 volumes of a saturated solution of harium

hydroxide

Basic Lead Acetate Solution. This solution possesses the following formula

Lead acetate	180 g
Lead oxide (litharge)	110 g
Distilled water to make	1000 g

Dissolve the lead acetate in about 700 ml distilled water, with boiling Add this hot solution to the finely powdered lead oxide and boil for onehalf hour with occasional stirring Cool, filter, and add sufficient distilled water to the filtrate to make the weight 1 kg

Benedict's Solution. Benedict s modification of the Fehling solution is stable even upon long standing. It has the following composition

Copper sulfate	17 3 g
Sodium citrate	173 0 g.
Sodium carbonate	100 0 g.
Distilled water to make	1 liter

With the aid of heat dissolve the sodium citrate and carbonate in about 800 ml water Pour (through a folded filter paper if necessary) into a class graduate and make up to 8.0 ml Dissolve the copper sulfate 10 about 100 ml water Pour the carbonate-citrate solution into a large beaker or casscrole, and add the copper sulfate solution slowly with constant stirring and make up to I liter The mixed solution is ready for use and does not deteriorate upon long standing

Benedict's Quantitative Sugar Reagent.

Copper sulfate (crystallized)		0 g	
Sodium carbonate (crystallized, one-half the weight of the			
anhy drous salt may be used)	200	0 g	
Sodium or potassium citrate	200	0 g	
Potassium thiocyanate	125	0 g	
Potassium ferrocyanide (a per cent solution)	5	0 ml	
Distilled water to make a total volume of	1000	0 ml	

With the aid of heat dissolve the earbonate, citrate, and thiocyanate iu enough water to make about 800 ml of the mixture and filter if necessary Dissolve the copper sulfate separately in about 100 ml water and

pour the solution slowly into the other liquid, with constant stirring Add the ferrocyanide solution, cool, and dilute to exactly 1 liter Of the various constituents, the conper salt only need be weighed with exactness Twenty five ml of the reagent are reduced by 50 mg of glucose

Benedict's Sulfur Reagent.

Crystallized copper mitrate, sulfur free or of known	sulfur
content	200 g
Sodium or potassium chlorate	50 g
Distilled water to	$1000 \mathrm{ml}$

Benzidine Solution. Place 4.33 ml glacial acetic acid in a small Erlenmeyer flask, warm to 50° C, and add 0.5 g of benzidine. Heat the flask for eight to ten minutes in water at 50° To the resultant solution add 19 ml of distilled water This solution may be kept for several days without deteriors tion

Bial's Readent.

Oremol	15g
Concentrated HCl	500 ml
Ferric chloride (10 per cent)	20-30 drops

α, α'-Bipyridine Solution Dissolve 0 2 g α, α' bipyridine (Edwal Laboratories, 732 Federal St , Chicago) in 100 ml 10 per cent acetic acid Biuret Paper (Kantor and Gles) Immerse filter paper in Gies'

Biuret Reagent (below), then dry and cut into strips

Bruret Reagent (Gies) This reagent consists of 10 per cent KOH solution to which 25 ml 3 per cent CuSO, solution per liter has been added This imparts a slight though distinct color to the clear liquid

Biuret Reagent (Welker) Add 1 per cent copper sulfate solution, drop by drop, with constant stirring to some 40 per cent sodium hydrovide solution until the mixture assumes a deep blue color

Black's Reagent. Make by dissolving 5 g ferric chloride and 0 1 g ferrous chloride in 100 ml water

Boneblack, Purification of. Treat 250 g commercial boneblack with 1000 ml 1 4 HCl solution Boil for half an hour Filter the boneblack from the acid solution by means of a Buchner funnel and aspirating pump Wash with hot water until the washings are neutral to litmus paper Dry and powder

Bromine Water. Add a few drops of hound bromine (Caulion. Very corrosue') to about 100 ml H2O, and shake Prepare fresh when reagent becomes colorless

Buffer Solutions. Standard. See no 35 to 37

Carmine-fibrin. Run fibrin through a meat chopper, wash carefully, and place in a 0 5 per cent ammoniacal carmine solution (1 g carmine dissolved in 1 ml ammonia and diluted to 200 ml) for 24 bours or until the maximum coloration of the fibrin (a dark red) is obtained The fibrin is then washed in water and water acidified with acetic acid It is preserved under glycerol

Cleaning Solution. To a few grams of sodium bicbromate crystals dissolved in a minimum quantity of water, add about 500 ml concen

trated sulfuric acid (technical grade is satisfactory)

Cochineal Solution. A saturated solution of cochineal in 30 per cent alcohol

Congo Red. Dissolve 0 5 g Congo red in 90 ml water and add 10 ml

95 per cent alcohol

Congo Red-Fibrin. This may be prepared by placing fibrin in faintly alkaline Congo red solution and heating to 80° C The fibrin is then washed and preserved under glycerol

Copper Sulfate Solutions for Specific Gravity Determination. A SATURATED COPPER SULPATE SOLUTION Place 4 lb "fine erystals," or pulverized CuSO, 5H2O, in a 1-liter bottle and add about 25 liters distilled water Stopper and shake vigorously for five minutes Immediately at the close of the sbaking period, insert a thermometer into the solution and record the temperature to the nearest half degree Centigrade When this has been done, immediately decant the supernaturt solution from the excess solid then clarify by filtration through cotton or a dry filter paper into a clean dry bottle This solution, which has been saturated with copper sulfate at a known temperature and 15 therefore of known composition is used at once to make up a stock copper

sulfate solution of ap gr 1 100

B STOCK COPIER SULFATE SOLUTION OF SP GR 1 100 From the table (p. 1325) determine the volume of saturated copper sulfate solution which is to be diluted to 1 liter to prepare a solution of sp gr 1 100 Measure out the indicated volume of saturated copper sulfate solution into a 500-inl graduated cylinder Pour the cylinder contents into a 1-liter volumetric flash allowing the cylinder to drain for 30 seconds. Dilute the flask contents to the mark with distilled water, stopper, and mix well by inversion. Because of a contraction in volume, the meniscus of the solution will now be somewhat below the mark. Allow to stand for one minute for drainage then add enough distilled water to bring the memseus again to the mark stopper and again mix well by inversion Pour into a clean dry 1-liter bottle Rinse the flask with distilled water, discarding the rinings, and repeat the above preparation three more time thus preparing I liters stock copper sulfate solution of 'p gr 100

This stock solution used for the preparation of the standards described below should be labeled with its temperature at the time of preparation since it may be used indefinitely provided the tempera-

VOLUME OF SATURABLE COLPUT SULFATE SOLUTION TO DILUTE TO 1 LITER TO PRELARE STOCK SOLUTION OF D₁₅ = 1 1000

Tempero the Sat Soluti the Ti Satur	uraled on at me of	Volume of Solution to Dilute to 1 Liter	the Sa Solut the T	olure of lurated ion at ime of olion	Volume of Solution to Dilute to 1 Liter	the So Solu the T	rature of sturated ston at sime of ration	Volume of Solution to Dilute to 1 Liter
° C	° F	ш	° C	° F	VI	° C	° F	WI
10 0	50 O	587	20 0	68 0	489	30 0	86 0	424
10 5	50.9	581	20 5	68 9	485	30 5	86 9	421
11 0	51 8	575	21 0	69 8	481	31 0	87.8	418
11 5	a2 7	569	21 5	70 7	477	31 5	88 7	415
12 0	53 6	563	22 0	71 6	474	32 0	89 6	412
12 5	54 5	557	22 5	72.5	470	32 5	90 5	410
13 0	5o 4	552	23 0	73 4	466	33 0	91 4	407
13 5	56 3	546	23 5	74 3	463	33 5	92 3	404
14 0	57 2	541	24 0	75 2	459	34 0	93 2	401
14 5	58 1	536	24 5	76 1	456	34 5	94 1	398
15 0	50 0	531	25 0	77 0	453	35 G	95 0	39 ₉
15 5	59 0	527	25 5	77 9	450	35 5	95 0	392
16 0	60.8	522	20 0	78.8	446	36 0	96 8	389
16 5	61 7	518	26 a	70 7	443	36 5	97 7	387
17 0	62 6	514	27 0	80 6	440	37 0	98 6	384
17 5	63 5	509	27 5	81 5	438	37 5	99 5	381
18 0	64 4	505	28 0	82 4	435	38 0	100 4	378
18 5	65 3	501	28 5	83 3	432	38 5	101 3	374
10 0	66 2	497	29 0	84 2	429	39 0	102 2	371
19 a	67 1	493	29 5	85 1	427	39 5	103 1	368
20 0	68 0	489	30 0	86 0	424	10 0	101 0	365

c Standard Copper Sulfate Solutions of Known Specific Grayter A 100 ml portion of copper sulfate solution of any desired sp gr between 1 008 and 1 075, accurate to ± 0 0003, may be prepared from the stock solution of sp gr 1 100 as follows measure from a burset into a 100-ml volumetric flask a volume of stock solution less by 1 ml than the value represented by the second and third decimal places of the desired sp gr (For instance, if the desired sp gr is 1 055, 54 ml of stock solution are taken, for a sp gr of 1 017, 16 ml are taken and so on) After measuring out the stock solution, dilute the contents of the 100-ml flask to the mark with distilled water and mix well. This solution may be used at any temperature within 15° to 20° C of the temperature at preparation Smaller portions than 100 ml may be prepared by using proportionately reduced volumes.

Cross and Bevan's Reagent. Prepare by combining 2 parts of concentrated hydrochlone acid and 1 part of zinc chloride, by weight

Digitonin Solution. Dissolve 1 g digitonin (Hoffmann-La Roche, Inc., Nutley, N J) in 1 liter of distilled water and place in a refrigerator for 24 bours. Filter Concentrate the entire filtrate and make up to ex-

actly 500 ml (To hasten evaporation draw a rapid stream of air, filtered through cotton, over the surface of the solution in a tared flass, which is immersed in a boiling water bath until the desired weight is reached.)

Duponol C (E I du Pont de \emours and Co, Wilmington, Dela

ware) Sodium salt of a sulfuric ester of lauryl alcohol

Ehrlich's Aldehyde Reagent A solution of p-dimethylaminobenzaldehyde acidified with HCl

Ehrlich's Diazo Reagent Two separate solutions should be prepared and mixed in definite proportions when needed for use

(a) 5 g sodium nitrite dissolved in 1 liter distilled water

(b) 5 g sulfamilie acid and 50 ml hydrochloric acid in 1 liter distilled water

Solutions (a) and (b) should be preserved in well stoppered vessels and mixed in the proportion 1-30 when required It is said that greater delicacy is secured by mixing the solutions in the proportion 1-100 \(\) satisfactory modification of this reagent containing less acid is described on p. 503. The sodium mitnic deteriorates upon standing and becomes unfit for use in the course of a few weeks.

Esbach's Reagent Dissolve 10 g pierie acid and 20 g citric acid in 1

hter water

Exton's Reagent Dissolve 200 g $\agnumber \agnumber \ag$

Fehling's Solution Fehling's solution is a mixture of copper sulfate

solution and alkaline tartrate solution prepared as follows

(opper sulfate solution = 34 65 g copper sulfate dissolved in water and made up to 500 ml. Waline tartrate solution = 125 g potassium hydroxide and 173 g Rochelle salt dissolved in water and made up to 500 ml. Günzberg's Reagent. Dissolve 2 g. phloroglucinol and 1 g. vauiilin in 100 ml, 95 per cent alcohol.

Halnes' Solution. This solution may be prepared by dissolving 8.314 g. copper sulfate in 400 ml. water, adding 40 ml. glyccrol and 500 ml. 5 per cent potassium hydroxide solution.

Hopkins-Cole Reagent. To 1 liter of a saturated solution of oxalic acid add 60 g. sodium amalgam and allow the mixture to stand until the evolution of gas ceases. Filter and dilute with 2 to 3 volumes of water.

Hopkins-Cole Reagent (Benedict's Modification). Place 10 g. powdered magnesium in a large Erlenmeyer flask and shake up with enough distilled water to liberally cover the magnesium. Now slowly add 250 ml. of a cold, saturated solution of oxalie acid. The reaction proceeds very rapidly and with the liberation of much beat, so that the flask should be cooled under running water during the addition of the acid. The contents of the flask are shaken after the addition of the last portion of the acid and then poured upon a filter, to remove the insoluble magnesium oxalate. A little wash water is poured through the filter, the filtrate acidified with acetic acid to prevent the partial precipitation of the magnesium on long standing, and made up to a liter with distilled water. This solution contains only the magnesium salt of glyovylic acid.

Hübl's Iodine Solution. A solution of iodine and mercuric chloride used to determine iodine numbers of unsaturated compounds. (See Iodine

Solution, Wijs.)

Hydrochloric Acid Standard (0.1 N) Solution. See p. 868.

Hypobromite Solution. The ingredients of this solution should be prepared in the form of two separate solutions which may be united as needed.

(a) Dissolve 125 g. sodium hromide in water, add 125 g. hromine, and make the total volume of the solution 1 liter.

(b) A solution of sodium hydroxide having a specific gravity of 1.25.

This is approximately a 22.5 per cent solution.

Preserve both solutions in rubher-stoppered hottles and when needed for use mix 1 volume of solution (a), 1 volume of solution (b), and 3 volumes of water.

Iodine Solution. Prepare a 2 per cent solution of potassium iodide

and add sufficient iodine to color it a deep yellow.

Iodine Solution (0.1 N). Weigh 12.685 g. pure resublimed iodine into a small weighing bottle using a porcelain spatula Dissolve 18 g. pure KI in ahout 150 ml. water. Transfer the iodine to a liter flask, washing out the last traces with some of the KI solution, which is then poured into the flask. Stopper and shake occasionally until dissolved, if necessary, a few more crystals of KI may be added to aid solution. Dilute to the mark and mix well. Keep in a glass-stoppered bottle in a cool dark place. Standardize at once against 0.1 N sodium thiosulfate solution. Measure out accurately 25 ml. of the iodine solution into an Erleimeyer flask, run m sodium thiosulfate until the color is pale yellow, then add a few ml. of a 1 per cent solution of starch (preferably soluble starch) and titrate to disappearance of blue color. Care should be taken near the end point.

Iodine Solution (Wijs). Weigh into a 300-ml. flask 9.4 g. iodine trichloride. Add about 200 ml. glacial acetie acid. Stoppor with a cork

carrying a CaCl₂ tube and heat on the water bath until solution is complete. Rub 7.2 g iodine to a fine powder in a mortar, wash with glacial acetic acid into a second flash, and heat this in the same way to dissolve the iodine. Pour the contents of hoth flasks into a liter volumetric flask. Add glacial acetic acid. Measure 10 ml. of the solution into a 500 ml. Erlenneyer flash. Add 10 ml. 10 per cent KI and about 200 ml. water litrate with standard sodium throsulfate solution and determine the iodine equivalent of 1 ml. of the solution.

Iodine-Zinc Chloride Reagent. Dissolve 20 g ZnCl; in 85 ml water and, when cool, introduce the iodine solution (3 g KI and 15 g In 60 ml water) drop by drop until todine begins to precipitate

Kraut's Reagent Dissolve 272 g KI in water and add 80 g bismuth subintrate dissolved in 200 g HNO, (sp gr 118) Permit the KNO to crystallize out, then filter it off and make the filtrate up to 1 liter with water

Lead Acetate, Basic. (See Basic Lead Acetate)

Lime Water. Shake up an excess of calcium oxido or hydroxide with distilled water and leave well stoppered over night Decant clear supernatant solution and keep free from CO. of ar

Litmus-Milk Powder. Add 1 part powdered litmus to 50 parts dried milk powder To make a litmus milk solution, add 1 part of this powder to 9 parts water

Lohmann's Reagent. Add 25 ml concentrated mirre acid to 100 g mercine nitrate octahydrate followed by 25 ml water Warm to dissolve Lugol's Solution. Dissolve 5 g todino and 10 g potassium jodide in

100 ml distilled water

Magnesia Mixture Dissolve 175 g magnesium sulfate and 350 g ammonium chloride in 160 ml distilled water Add 700 g concentrated ammonium hydroxide mix thoroughly, and preserve the mixturo in a klass-storopered bottle

Magnesium Nitrate Solution for Ignition Dissolve 320 g calcined magnesia in nitra acid, avoiding an excess of the latter then add a little calcined magnesia in excess boil filter from the excess of magnesia ferrie oxide, etc., and dilute with water to 2 liters

Methyl Orange Dissolve 01 g methyl orange in 100 ml distilled

 α Naphthol Solution. Dissolve 1 g α -naphthol m 100 ml $\,95$ per eent aleohol

Nessler's Reagent: (1) Formula of Folin and Wu Nessler's solution is an alkaline solution of the double iodide of mercury and potassium (HgI22KI) Into a 500-ml Florence flask introduce 150 g potassium iodide and 100 g iodine, add 100 ml water and an excess of metallic mercury, 140 to 150 g Shake the flask continuously and vigorously for 7 to 15 minutes or until the dissolved iodine has nearly all disappeared Tbe solution becomes quite hot When the red iodine solution begins to become visibly pale, though still red, cool in running water and continue the shaking until the reddish color of the iodine has been replaced by the greenish color of the double iodide. The whole operation does not usually take more than 15 minutes. Decant the solution, washing mercury and flask with liberal quantities of distilled water. Dilute the solution and washings to 2 liters. If the cooling was be, in in time, the resulting reagent is clear enough for immediate dilution with 10 per ceut alkali and water and the finished solution can be used at once for nesslerization.

From this stock solution of potassium mercuric iodide prepare final Nessler's solution as follows. Into a flask of at least 5 liters capacity introduce 3500 ml. 10 per cent sodium hydroxide solutiou, add 750 ml. of the double iodide solution and 750 ml. distribled water, making a liters of solution. The 10 per cent NaOH should be made from a saturated solution (containing about 75 g. per 100 ml.) which has been allowed to stand until the carbonate has settled, the elear solution being decanted and used. This solution should have been standardized with an accuracy of at least 5 per cent. Nessler's reagent should be used in the ratio of 10 ml. per 100 ml. of solution to be nesslerized, except where excessive amounts of ands are prescut, as in direct nesslerization procedures.

The alkalinity of the Nessler's reagent is important and may be checked by titrating with it 20 ml portions of N HCl A good end point with phenolphthalein should be obtained at 11 to 11 5 ml If as httle as 9 5 ml is required, the solution is too alkaline One ml of the dilute (I I) and digestion mixture should also require 9 to 93 ml Nessler's solution to

neutralize it

(a) Formula of Roch and McMeerin Dissolve 22 5 g todine in water containing 30 g potassium onded After the solution is complete, add 30 g put metallie mercury, and shake the mixture well, keeping it from becoming hot by immersing it in tap water from time to time. Continue this until the supernatant haud has lost all of the yellow color due to todine. Decant the supernatant aqueous solution and test a portion by adding a few drops thereof to 1 ml of a 1 per cent soluble starch solution. Unless the starch test for ordine is obtained the solution may contain mercurous compounds. To the remaining solution add a few drops of an iodine solution of the same concentration as employed above, until a faint excess of free iodine can be detected by adding a few drops thereof to 1 ml of the starch solution. Dilute to 200 ml and mix well. To 975 ml of an accurately prepared 10 per cent sodium by droude solution now add the entire solution of potassium mercuric iodide prepared above. Mix thoroughly and allow to clear by standing

(c) FORWILL OF BOCK AND BENEDICT Place 100 g mercuric iodide and 70 g potrassium iodide in a liter volumetric flask and add about 400 ml water Rotate until solution is complete Now dissolve 100 g

NaOH in about 500 ml water, cool thoroughly and add with constant shaking to the mixture in the flask, then make up with water to the liter mark. This usually becomes perfectly clear When the small amount of brownish red precipitate which forms settles out, the supernatant fluid is ready to be poured off and used

Neutral Olive Oil. Shake ordinary olive oil with a 10 per cent solution of sodium carbonate, extract the mixture with ether, and remove the

ether by evaporation. The residue is neutral olive oil

Neutral Red. A 1 per cent solution in 50 per cent alcohol

p-Nitrophenol. A I per cent solution in a0 per cent alcohol

Nylander's Reagent. Digest 2g bismuth subnitrate and 4 g Rochelle salt in 100 ml of a 10 per cent solution of potassium hydroxide The reagent should then he cooled and filtered

Ohermayer's Reagent. Add 2 to 4 g of ferric chloride to a liter of

hydrochloric acid (sp gr 1 19)

Oxalic Acid Standard (0.1 N) Solution. See p 867

Permutit. A synthetic aluminum silicate ohtained from the Permutit Company, New York. Only such preparations as pass through a 60-mesh sieve but not through an 80-mesh sieve should be used It should give off very little dust or turbid material to water and settle in a few seconds It may be used more than once hy washing first with water, then with 2 per cent acetie acid and finally with water again

Phenol Reagent (Folin and Ciocalteu). See n 939

Phenolphthalein. Dissolve 1 g phenolphthalein in 100 ml 95 per

cent alcohol

Phenylhydrazine Acetate Solution. This solution is prepared by mixing 1 volume glacial acetic acid, 1 volume water, and 2 volumes

plicas lhy drazine (the hase) Phenyihydrazine Mixture. This mixture is prepared by combining 2 parts phenylhydrazine hydrochloride and 3 parts sodium acetate bj weight These are thoroughly mixed in a mortar A mixture of better keeping quality may be made by mixing equal weights of the hydro-

chloride and anhydrous sodium acetate Phosphomolyhdic Acid. Saturate some sodium carbonate solution with pure molybdic acid Add 1 part crystalline disodium phosphate to 5 parts acid and evaporate to dryness Fuse in a porcelain dish at a dull red heat Dissolve the sodium phosphomolybdate in 10 parts water and

add nitric acid until the solution turns a golden vellow color

Phosphotungstic Acid. Dissolve 100 g sodium tungstate and 60 to 80 or disodium nhosphate in 500 ml water. Add nitric acid to an acid watch glass and let stand over night at room temperature. If erystallization fails to occur, seed with a small crystal of pure pierie acid. At the end of two hours, filter with suction on a hardened filter, and wash with about 35 ml. cold glacial acetic acid. Suck as free from acetic acid as possible and dry at about 80° to 90° C, with occasional stirring, until there is no odor of acetic acid. Conduct operations in a good current of air.

B. SODIUM PICRATE METHOD OF FOLIN. Transfer 500 g. moist pierie acid to a Florence flask of 1500 ml. capacity. Add 500 ml. acetone. Shake, with a little warming under hot tap water, until all the crystals have dissolved. Add 20 g. active charecal ("Norit"). Shake, and filter into another flask. During this filtration keep the funnel closed with a watch

glass to prevent evaporation.

Dissolve 250 g. anhydrous sodium carbonate and 100 g. sodium chloride in 2500 ml. warm water in a 4-liter beaker. While stirring with an agateware spoon, add the acctone solution gradually to the alkaliue salt solution. When the reaction (CO₂ evolution) is finished, let stand, preferably in cold water, for about 30 minutes, and filter on a large Buchner funnel (diam 20 cm.). Wash with about 2 liters sodium chloride solution (7 per cent) and suck as dry as possible.

If the original picric acid is of good quality, the sodium picrate on the Buchner funnel will be pure, but it is a little safer to recrystallize it once

as follows:

Return the precipitate to the 4-liter beaker and add 2 liters holling water and 20 g. sodium earhonate. To the resulting hot solution add gradually, with stirring, 150 g. sodium ehloride, cool, filter, and wash as before with 7 per cent sodium ehloride solution. Then wash once or twice with a more dilute sodium ehloride solution (2 per cent) and finally wash once with methyl alcohol to remove most of the remaining chloride and water. Dry, either at room temperature or over a radiator.

Test for the Puntty of Sobium Pichate. Make 100 ml. of a 3 per cent solution. Transfer 5 ml. and 10 ml. to test tubes graduated at 25 ml. Dilute each to about 22 ml., add 2 ml. 5 per cent sodium hydroxide, dilute to volume, mix, and let stand for 10 minutes. Then add 4 g. powdered potassium chloride, mix hy inversion for about 1 minute, filter on a 9-cm quantitative filter paper, and compare the two filtrates in the colorimeter. If the picrate is pure, the two filtrates will have the same color.

Picric Acid. The process for the preparation of pure picric acid is exactly the same as described above, up to the final washing with methyl alcohol, except that hardened filter paper should he used on the Buchner funnel.

One simply converts the purified sodium picrate in the Buchner funnel into picrie acid by treating it with dilute hydrochloric acid.

Prepare at least 2 liters of such acid (1 volume conc. acid to 4 volumes water). Disconnect the filtering flask from the suction pump. Pour the acid over the pierate. Stir up the precipitate with a porcelain or glass spoon, so as to make sure that the acid has acted on it all. Use plenty of the acid. Unchanged pierate can he distinguished by its darker color. When no more pierate is visible, connect again with the suction pump, and filter to dryness. Then wash five or six times with cold distilled water and suck as dry as possible. Temperatures up to 90° C. can safely be used for the drying of pieric acid.

Pictic Acid. Saturated Solution. This may be prepared either by allowing distilled water to stand in contact with an excess of pieric acid with occasional shaking or by making a 12 per cent solution

Potassium Mercuric Iodide Dissolve 6775 g dry crystalline mercuric chloride and 25 g potassium iodide separately in water Mix.

Dulute to 1000 ml

Potassium Permanganate Standard (0 1 N) Solution. Dissolve 3 162 g pure notassium permanganate in a liter of distilled water, allow to stand a few days, and filter through glass wool Standardize against 0 I \ oxalic acid solution or against pure dry sodium or potassum ovalate One ml 01 \ permanganate is equivalent to 67 mg sodum oxalate (See also footnote 200. p 645)

Potassium Persulfate Solution (Saturated). Add 100 ml distilled water to 7 g pure potassium persulfate in a glass-stoppered bottle

Undissolved excess settles and compensates for loss by decomposition. Ringer's Solution To 960 ml 0 104 M \aCl solution add 20 ml 0 L54 M KCl solution and 20 ml 0 11 M CaCl, solution

Roberts' Reagent Mix I volume concentrated nitric acid and

5 volumes saturated solution of magnesium sulfate

Rosenheim's Iodo-potassium Iodide Solution Dissolve 2 g rodine and 6 g potassium rodide in 100 ml water (This is different from I ugol s solution in which the proportion of iodine to notassium iodide 18 Ĭ 2)

Sahll's Reagent This reagent consists of a mixture of equal parts of 48 per cent solution of potassium iodide and 8 per cent solution of

notassium iodate

Schweitzer's Reagent To 10 parts ammonium hydroxide (sp gr 0 90) add 3 parts distilled water To this mixture add a slight excess of copper carbonate shake vigorously and allow to stand over night Suplon off the clear, supernatant hound

Selivanoff's Reagent Dissolve 000 g resorcinol in 100 ml dilute

(1 2) hydrochloric acid

Sodium Acetate Solution Dissolve 100 g sodium acetate in 800 ml d stilled water, add 100 ml 30 per cent acctic acid to the solution, and make the volume of the mixture up to 1 liter with distilled water

Sodium Mcoholate (0 1 N) Solution The sodium alcoholate 1 made by dis olving 23 g cleaned metallic sodium in 1 liter absolute declied It may be standardized against pure henzoic acid in washed chleroform or against 0.1 \ HCl provided the alcoholate solution con tams not mere than traces of carbonate

Sodium Alizarin Sulfonate Dissolve 1 g sodium alizarin sul

fonate in 100 inf. water

Sodium Cobaltinitrite Solution Prepare according to Kramer and lusdall as follo vs

the gas has passed off The reagent is placed in the ice chest and filtered each time before using It will keep at least one month

Fo 20 ml of this sodium cobaltinitrite solution add 2 ml 40 per cent silver intrate solution. Shake vigorously and filter to remove trace of

insoluble precipitate

Sodium Hydroxide (Saturated Solution). Shake up about 110 g of best quality NaOH with 100 ml distilled water in a 300-ml Erlenmey er flask ("Pyrex") Stopper and allow to stand for a couple of days or until the sodium carbonate settles to the bottom leaving a clear solution of NaOH practically free from carbonate, and containing about 75 g NaOH per 100 ml

Sodium Hydroxide Standard (0.1 N) Solution. Sec pp 867 and 868

Sodium Thiosulfate Standard (0.1 N) Solution. Weigh out 25 g ordinary e p sodium thiosulfate or 24 83 g of the pure dry recrystallized salt Dissolve in water and dilute to a liter Boiled distilled water must be used Keep in a bottle with a siphon arrangement and carrying a sodalime tube to evelude CO. It is best standardized against acid potassium iodate KH(IO₃). Weigh out accurately 0 3249 g acid potassium iodate Dissolve in off maker, heating gently if necessary Transfer the solution to a 100 ml flash, rinsing the beaker earefully and make to mark with water This solution is exactly decinormal Pipet 25 ml into an Erleimeyer flash, add 1 g potassium iodide dissolved in a little water, and a few cubic centimeters of dilute hydrochloric acid Titrate immediately with the thiosulfate solution. When the solution becomes pale yellow add a few ml 1 per cent solution of soluble starch and titrate to loss of filiu color.

Sodium Tungstate Solution. A 10 per cent solution of sodium tungstate in water The ep sodium tungstate made by the J T Baker Chemical Co, Phillipsburg, N J, or by the Mallinckrodt Chemical Co, St Louis, No, is satisfactory (See also p 551) If not easily soluble, prepare a hot 10 per cent solution, cool, and titrate 25 ml with 10 per cent NaOH and phenolphthalein to a pink color which lasts for 3 minutes

Add a proportional amount of NaOH to the main solution

Soluble Starch Suspend 3 g raw potato starch in 100 ml redistilled 95 per cent alcohol, add 0 75 ml concentrated HCl, and heat on a boiling water bath with a reflux condenser for exactly 10 minutes Add at once an amount of normal sodium bicarbonate solution just sufficient to neutral 12e 0 75 ml of the acid, using methyl orange as indicator Decant through a filter and wash with several portions of alcohol Dry at room temperature, seve and preserve

Soluble Starch Solution A solution of soluble starch suitable for most nodometric tirations and with good keeping qualities is made according to Pincussen by dissolving 1 g soluble starch in 10 ml boiling

water and adding to 90 ml saturated NaCl solution

Starch-Iodic Acid Test Paper This test paper is prepared as follows Saturate a good quality of filter paper with 0 per cent starch paste to which has been added sufficient todic acid to make 1 1 per cent solution of iodic acid and allow the paper to dry in the air Cut it in strips of suitable size and preserve for use

Starch Paste. Grind 2 g stareh powder in a mortar with a small amount of water Bring 200 ml water to the boiling point and add the starch mixture from the mortar with continuous stirring Bring again to the boiling point and allow it to cool This makes an approximate 1 per cent starch paste which is a very satisfactory strength for general use

Stoke's Reagent. A solution containing 2 per cent ferrous sulfate and 3 per cent tartanc and When needed for use a small amount should be placed in a test tube and ammonium hydroxide added until the preputate which forms on the first addition of the hydroxide has entirely dissolved This produces ammonium ferrolartrate, which is a reducing

agent
Sulfuric Acid, N/12. Add 2.5 ml concentrated sulfuric acid to a
liter of distilled water Standardize against alkali of known strength
(See p. 544)

Sulfuric Acid, Two-thirds Normal. Add 35 g concentrated cp sulfunc acid to a liter of distilled water Standardize against alkalı of

known strength
Takayama's Solution. A mixture of 3 ml 10 per cent NaOH, 3 ml
pyndine, 3 ml saturated solution of gifteosy and,7 ml water The solution works rapidly in the cold if at jeast 21 hours old With a fresh solution, warming or more time is necessary. It keeps for one to two months

Tannic Acid A freshly prepared per cent aqueous solution

Tanret's Reagent Dissolve 135 g mercuric chlorido in 25 ml water, add to this solution 3 32 g potassium iodide dissolved in 25 ml water, then make the total solution up to 60 ml with distilled water and add 20 ml clacial acetic acid to the mixture

Therefore Toldine Dissolve 70 g todane and 50 g potassium todide in 50 ml distilled water Dilute to 1000 ml with 95 per cent alcohol

Topfer's Reagent. Dissolve 0.5 g dimethylaminoazobenzene in

Tropeolin 00. Dissolvo 0 05 g tropeolin 00 m 100 ml 50 per cent

Tungstate-Molybdate Reagent To 10 g pure NH-free, molybdre acid in a flask add 50 ml N sodum hydroxide and boil gently for 3 to 5 minutes Add about 150 ml water Filter the hot solution and add to the filtrate a solution of 80 g sodium tungstate dissolved in about 600 ml water Dilute to 1000 ml

Uranium Acetate Solution. Dissolve about 35 0 g uranium acetate in 1 liter water with the aid of heat and 3 to 4 ml glacial acetic acid Let stand a few days and filter Standardize against a phosphate solution containing 0 000 g. P.O. per ml For this purpose dissolve 14 721 g pure air

y sodium ammonium phosphate (Na MidHPO4 + 4H₂O) in water to ake a liter To 20 ml of this phosphate solution in a 200-ml beaker add J ml water and J ml sodium acctate solution (see p 1332) and itirate ith the uranium solution to the correct end reaction as indicated in the ithod proper p 952. Il exactly 20 ml uranium solution are required ml of the solution is equivalent to 0.005 g P₂O₂. If stronger than this dute accordingly and check again by thration

Winkler's Reagent. Cuprous chloride 10 g ammonium chloride 0 g, distilled water to 170 ml. For use mix this solution with ammonium

ndroxide (sp gr 09) in the proportion of 3 1

II. COMMON ACIDS AND ALKALIES AS PURCHASED

Substance	Specific Gravity (at Room Temperature)	Per Cent by Weist	1 pproximate Normality
/II,0II	0 90	28	15
	1	(as \ H ₂)	
\aOH (saturated)	15	50	19
H SO ₄ (concentrated)	1 84	95	36
II \O. (concentrated)	1 42	70	16
IICl (concentrated)	1 19	36	12
II.PO.	171	85	45
(syrupy) leetic acid (glacial)	1 05	99 5	17

III. TABLE OF COMPOSITION OF FOODS*—RAW, PROCESSED,

SIGNS AND SYMBOLS USED

An asterisk in the table stub indicates an item for which the composi-

Parentheses denote imputed values for which little or no experimental evidence was available, for which there was relatively little basis for inputing a value from another form of the food, or for which reported data were not considered suitable. A zero in parenthesis is used where actual data were lacking and the amount of a constituent present was regarded as none or probably too little to measure.

Dashes show that no basis could be found for imputing a value although there was some reason to believe that a measurable amount of the constituent might be present

The word "Trace" is used to indicate vitamin values that would round to zero with the number of decimal places carried in these tables For other components that would round to zero, a zero is used A zero followed by a decimal point indicates that there may be up to 0.5 of the unit present but bases for showing the amount were inadequate Numbers with or without decimal points indicate that the average has been rounded to the nearest whole number or in the case of vitamin A to the nearest multiple of ten.

Abridged from Watt Merrill Orr Wu and Pecot Agriculture Handbook No 8
 Washington D.C. U.S. Department of Agriculture, 1950

In preparing this abridgment the following items have been omitted. Raw foods not commonly eaten certain canned foods when values for corresponding cooked items are included forces foods and prepared infant foods similar in composition to forms included condensed soups, lean or fat meats when medium fat meats are listed, cross references humbers missage from first column recression control uses.

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True than the rivedam and main are based on it en minima level of engelment apecifed in standards of identity proposed by it o Federal Security Agency and 1 Wf en the sugarst of nonfat mulk solule m con n erecal bread is unknown use bread with 4 per cent nonfat mulk solule stom 135 for unenrelied bread and 139 for published in the Federal Register August 3 1913

of the fact went in the recipe to butter or fortered in arganne the vitemin A value per 100 g. would be 540 [U in foundation cake, 430 I U in foundation cake food, 410 I U in dark Ingj cake 370 I U in plain cake 280 I U in plain cake seed 1900 I U in pound cake 130 I U in rich cake ned 000 I U in rich cake eved at it a calcium contributed by chocolate is considered unavailable the value would be 38 mg per 100 g * Year round average

Mytau in A based on deeply colored varieties

13 Based on products ranging from 2.5 to 6.6 mg per 100 g cereal. The nasen value of some products is as high as 23.6 mg " Calcium may not be available because of prozence of exalle acid

diosition of Foods, 100 G, Edista Pontion-(Continued)

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14460 | g if the added emulaifying agent does not contain phosphorus

11 Approximately one-that of that total amount of entrobydrate calendated by difference w starch and sugar. The remaning portion is made up of n akenals thought 14 Not less tign 60 per cent meat not more than 8 per cent cereals geasonings 16 Vitan in values are based on muscle meat only

19 Vitan in A lased on yellow corn white corn contains only a trace 12 Calcium nay not be available because of presence of exalte agid to be utilized only poorly if at all by the body

19 Dased on recipe using white corn meal if yellow corn meal is used the vatamin A value is 330 I U 11 Blaned on recipo using white corn heal if yellow corn meal is used the situmen A walue in 250 I U

19 Vitan in A based on yellow corn flour white corn Cour contains only a trace 28 Vitan in A based on yellow corn grits white corn grits contain only & trace

metic Act

11 Iron tlanmae aboffayn and maen are based on the maganum level of enorthment specified to shandards of identity promulgated under the Food Drug and Cos

11 Vitan in A based on yellow corn meal white corn meal contains only a trace

Constraction or Loods, 100 G., I purk Fourior—(Continued)

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ol cucumbers unpared contan about 12 mg 1700 and 200 I W valamun A per 100 g. s. nbofavna and nacm are based on the minimum levels of ennehment specioed na standards of identity promulgnted under tho Food Drug, and C	w Hens are from the medium fat wholesale cuts considered to be nearest approximations for sadiented retail Hems
11 Based on pared cucumbers u	netic Act ** Values for raw itenis are fron

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Ploy them ne riboflavin and macin are based on the minimum level of emochment specified in the standard, of identity promulgated under the Fo. d. Drug and 16 The vitanin values are based on the drained solids Cosmette Act

11 Based on the average vitamia A content of fortified marganne Most of the margannes manufactured for use in the United States lawe 15 000 I U of vitamin A

edded per pound. The innumum Federal apecifications for fortified margarine require the addition of 6 000 I U of vitamin A per pound

" Based on unfortified products 13 Total sugars

"Suinted ash Tius overestimates the ash in the range of S to 20 per cent

11 from this suine relociaves and much are based on the manneum towel of enrehment specified in the standards of admirty promulgated under the Dood. Drug and Coemetle Ant

Courosition of Foods, 100 G, Loine Portion—(Confinued)

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#TI a vitamin A values range from about 16 I U per 100 g of white fleahed plantains to 1 206 I U per 106 g of deeper yellow fleshed varieties * Values for raw items are from the medium fat wholesale cuts coasidered to be nearest approximations for indicated retail iter s "The proximate constituents calcium phosphorus and vitamin A are calculated from a recipe "Calcum may not be available because of presence of oxaire acid

Couloming of 1 0008, 100 G, Points Portion-(Continued)

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41ron thanno riboflaviu and nincin are based on the minimum level of enrichment specified in the standards of identity of breade proposed by the Pederal So

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4 Year round average. Recently dug potatoes contain about 24 mg, ascorbic acid per 100 g. The value is only half as high after three months of storage and about one 41 Calclum may not be available because of presence of exalte acid third as high when potatoes have been stored as long as six months

#11 bones are discarded calcium content would be much lower Bones equal about 2 per cent of total contents of can curity Agency and published in the Federal Register. August 3, 1943. "Minerals and vitamins are calculated from a recipe

PORTION-(Continued) Ė ţ

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4 All the rendy to-serves out a are calculated from equal weights of the condensed soup and water except cream boup which was based on equal weights of the condenses soup and rulk

44 Navy bean meal with farnasceous flour up to 15 per cent

4 from thismme riboflavin and miscus are based on the minimum level of sarrehment specified in the standards of identity promulgated under the Food Drug and 4 Approxic ately 40 per cent of this total amount of carbohydrate calculated by difference is sugar atarch and dextrin. The remaining portion is made up of materials " Pea meal with farmaceous flour up to 15 per cent thought to be utilized only poorly if at all by the body

11 Calcium and phosphorus are based on dark brown sugar values would be lower for light brown sugar * If very pale vancties only were used the vitamin A value would be very much tower " Calcium may not be available because of presence of oxalic acid

Cosmette Act

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ii Values for raw itoms are from the medium fat wholesale cuts cansidered to be nearest approximations for indicated fetail items 11 Vitan in A value of tortilias made from yellow corn tortillas made from white corn have no vitamu A value

19 Data assume cut to be prepared by brasing or pol-roasting. Use af proportanate quantity of drippings would add opproximately 50 per cent more thismine and ma 11 Use of proportionate quantity of liquid would double annunt of thismne and macin and add une third mure ribuflavin and 25 per cent more riboflavin

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11 from thasmine eiboflavis and miscin are based on the miscinam lavel of ourchment apsended in the atandards of identity pronulgated under the Frod Drug and of Figures for moisture are based on product as it reaches the mill prior to tempering ather prexis are constituents are adjusted to this basis

Ploo thisnies riboflavin and macin are based on thin minimm lovel of enrichment apenfied in the standards of identity promulgated under the Food Drug and Commete Act Calcium is based on the level usually found in self rissing flour which is in excess of the minimum (500 ing per pound) required Connette Act

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IV. AVERAGE PORTIONS OF FOODS Caloric Values and Acid- and Base-Forming Effects

The data presented here are reproduced by permission from the 1946 Nutritional Charts prepared by the H. J. Heinz Co., Pittshurgh, Pa. The alkaline or acid effect is given in terms of ml. of N alkali (+) or acid (-) corresponding to 100 g. or 100 ml. of food (see discussion on p. 1097). The measure of "average portion" as given is hased upon the edible portion only and upon articles of average size. The food is calculated as in the raw state. These figures are not offered as exact data but merely as a guide in estimating average portions or servings; naturally there is considerable possible variation in this respect. For precise nutritional studies, all food consumed should be accurately measured. In these tables, T. = table-spoon, t. = teaspoon, e. = cup.

VEGETABLES

	1 2012	1,000,00		
	Alkaline		Average Portion	
Name	(+) 67 leid () Effect	Total Cal- ortes	Measure	16 eight
				grams
Artichokes (globe)	+76	50	1 heart, edible leaf portion	100
Arparagus	+08	23	6 stalks	ן זעט
Bamboo shoots	+80	30	24 c	100
Beans, baked	+40	59	14 c	100
Beans, dried	-18 0	94	2 T shelled	23
D	+54	37	I e stringless	100
Beans, green	+18 0	88	24 c shelled	100
Beans, green kidney Beans, dried lima	+42 0	91	16 c shelled	23
Beans, green lima	+28 0	125	24 e shelled	100
Beans, green uma Beans, soy, dried	+45	119	3 T shelled	28
		l	34 c shelled	100
Beans, soy, green	+17 0	158		100
Beans, soy, sprouts	+16 0	66	1 c	100
Beets	+11 0 +27 0	42	35 c	100
Beet greens Broccoli	+ 9 3	28 32	le "eurd"	100
	1	1		100
Brus-els aprouts	+11 0	53	6	57
Cabbage	+60		34 c shredded as slaw	100
Carrots	+11 0	1 40	1 large, scraped	100
Cauliflower	+ 53	27	le "curd"	100
Celerac (celery root)	+ 88	39	% c pared	100
Celery	+78	8	2 stalks	40
Chard, leaves	+160	22	11½ c	100
Chives	+13 0	52	2 bunches	100
Collarda	+	4.5	34 c	
Corn, canned (yellow)	- 18	84	⅓ c	115
Corn, green (yellow)	- 2 0			100
Cucumbers	+ 79			57
Dandelion greens	+18 0]1 c	100
Eggplant	+ 6 1			100
Endive	+ 7 6) 9	1/2 head	45
Escarole (chicory)	+40) 3		16
Garlic		.	1 clove, peried	10
Horseraduh	1 + 4 8	3 9	11:	10

APPENDIX IV

VEGETABLES-(Continued)

	Maline		Average Portion	
N ame	(+) or leid (-) Effect	Total Cal- ories	Measure	Weight
hale	+ 7 7 + 8 0	45 32	1 c leves	grams 100 100
Kohlrabi Lambsquarters	780	44	1 c leaves	100
Leeks	+70	23	2 stalks	57
Lentils, dried	-16 0	91	2 T	28
Lettuce	+74	12	34 bead	75
Marrow, vegetable	+ 19	17	5 € C	100
Mushrooms Mustard greens Okra	+ 4 0 + 4 5	30 25 17 23	7 1 c leaves 5 pods	100 100 50 50
Onions	+ 15	23	1 sprig	i
Paraley	1		1 sprig	_
Parsmps Peas, dried Peas, green Peppers, green Potatoes, sweet	+12 0 + 5 0 + 1 3 + 1 7 + 6 7	75 92 92 92 24 175	1/2 large, scraped 2 T 3/4 c shelled 1 empty pod 1 pared	100 28 100 100 145
Points - L.	+70	101	1 pared	120
Potatoes, white Pumpkins	+15	31	12 c seeded, rand removed	100
Radishes	+ 29	7	5	35
Rhubarb	+85	15	1 c stems	100 100
Rutabagas	+85	36	34 c scraped	100
Salsify (oyster plant) Sauerkraut	+ 2 9 + 5 7	78 14 22	2 scraped 24 c 1 c leaves	100 100 100
Spinaeh	+27 0 + 1 0	17	1 c seeded, rind removed	100
Squash, summer Squash, winter	+ 26	38	I c seeded, and removed	100
oquasii, winter	1			100
Taro	+18 0	93	1 corm, pared	100
Tomatoes	+56	20	1 small, cored	100
Turnips	+ 27	30 32	1 c leaves	100
Turnip greens	+ 2 3	21	216 c leaves	100
Water eress	+12 0	1		
Yams Yautia, yellow	+ +15 0	155 110	1 tuber, pared 1 corm, pared	150 100

FRUITS

Name	١,	Laline					
	Ac	+) or id (—) Effect	Total Cal	-		Measure	Weight
	-			-			grams
	1			o	. 1	large, cored	150
Apples		+ 3 7				halves, stoned	100
Apricots		+61		4		pear, pared, stoned	100
Avocados		+11 0	25			small, peeled	100
Bananas		+ 5 6		6			100
Blackberries	1.	+ 69	1 4	16	1	c	
Blueberries (huckleberries)	1.	+ 27	1 6	33	34	í c	100
Cherries		+ 4 5	1	67		Satoned	100
Cranberries	1	_		48	lï	c	100
Currants		+63		48		c	100
	- 1	+11 0		92		stoned	30
Dates, dried	l	-	1		ľ		30
Figs, dried	l	+33 0	1	83		1/2	100
Gooseberries	- 1	+ 33	1	37	13	ું c	100
Grapefruit	1	+70	1	43	1	≨ c juice	
Grapes	l	+40	Į.	72	lı	bunch, seeded	100
Guavas	- 1	+80	1	5G		pared seeded	100
	- 1		Ţ		1.		100
Lemons	- 1	+ 50	ļ	40		2 c juice	100
Limes	- 1	+10 0	l	52		2 c juice	100
Loganberries	- 1	+73	Ţ	64		l c	100
Mangoes	- 1	+ 50	-	69	-14	½ pared, seeded	
Melon-Cantaloupes	- 1	+75	- 1	40	1	1/2 seeded, rind removed	200
Honeydew	- 1	,	1	68	-1	% seeded, rind removed	200
Muskmelons	ļ	+75	. 1	52	-1	1/2 seeded, rand removed	200
Watermelons	- 1	+ 27		58		1 slice, seeded	200
\ectarnes	i	+ 6 2		65		2 pared, stoned	100
	- 1				- 1	• • • • • • • • • • • • • • • • • • • •	20
Olives, green		+ :		35		5 small, stoned	100
Oranges		+50	>	48		1/2 c juice	
Papayas			. 1	40		12 seeded, rind removed	100
Peaches (yellow)		+ 5 !		49		1 large, pared, stoned	100
Tears		1 + *	- 1	64	٠,	2 halves, cored pared	
Persimmons		1	- 1	135	5	1 small, seeded	100
Pineapples		+6	8	57	7	2 slices, canned	100
Plums		-	† l	5	4	3 stoned	100
Pomegranates		+ 3	5	7	4	½ seeded	100
Prunes, dried		-	1	10	8	4 stewed, stoned	3
Quinces		+ 4	. 1	5		1 boiled as a source	10
Itaisins		+34		13		1 boiled, as a sauce	4
Rasphernes black		+ 3		6		% c seeded and seedies	~ 10
Raspberries red		+ 6			6	1/8 ℃ 1/4 ℃	10
Strawberries		+ 5			iG.	12 hulled	10
Tangerines		+ 5			IG.	2 peeled, seeded	10

APPENDIX IV

CLREALS AND BAKERY PRODUCTS

	Alkalıne		Average Portion	
A ame	(+) or Acid (-) Effect	Total Cal- ories	Vicasure	Weigh
				grams
Barley, pearled	-10 0	107	3 T	30
Bread, enriched white	- 40	130	2 slices	50
Bread, rye	- 5 2	125	3 slices	50
Bread, white	- 40	130	2 shces	50
Bread, whole wheat	- 3 6	129	2 slices	50
Cake, devils food, iced	- 63	202	1 piece	57
Cake, sponge	- 9 7	108	1 piece	37
Cookies (average)	- or +	106	1-2	28
Commeal, yellow	- 49	107	3 T	30
Crackers (Graham)	- 8 5	125	3	30
Crackers, soda	- 90	104	4	25
Doughnuts	- 17	242	1	57
Farma	-11 0	110	3 T	30
Farms, enriched	-11 0	110	3 T	30
Flour, buckwheat	- 7 1	77	2 T	22
Flour, rye	-11 0	64	2 T	18
Flour, soy bean	+95	82	2 T	18
Flour, white	- 90	60	2 T	17
Flour, whole wheat	-11 0	57	2 T	10 50
Hommy, white	- 17	178	14 c	1
Macaroni or spaghetti	-14 0	101	14 c	28 100
\oodles, egg	l –	382	34 c	25
Oatmeal (Rolled Oats)	-12 0	98	14 c	17
Popcorn, nonned	- 80	69	1 c	25
Pretzels	- 70	90	6	
Rice, brown	- 57	106	3 T	30
Rice, white	- 90	98	2 T	28
Tapioca	0.0	140	1/4 c	40 28
Wheat bran	-2a 0	87	1 c	28
Wheat germ	-20 0	76	2 T	
Wheat, whole	-12 0	72	2 T	20

grams 28 4 slices 149 - 48 113 144 14 lb -21 0 113 303

acon (lean) eef hrains Ив leef, chuck -11 0 113 325 eef, corned (medium) -10 0 113 1/4 lb Sect heart (lean) - 9 1 118

MEATS, FISH, AND POULTRY PRODUCTS-(Continued)

	T					Average Portion	
Name	(-	kaline +) or id (—) Effect	Co	tal al- tes		V easure	Weight
			_		_		grams
	1	- 8 5	١,	55	3 (113
Beef kidney		- 8 5 -11 0		49		lb.	113
Beef liver		-11 0		85		1b	113
Beef, loin		-11 0		263		1b	113
Beef steak		-12 0		163	lî		113
Beef sweetbreads	- 1	12 0	(1		75
D. f.t.		-11 0	1	170	15	slices	
Beef tongue Bluefisb	- 1	- 90		133		fish filet	113
Bologna	- 1	- 8 5	1	93	4	slices, skin removed	113
Chicken	- 1	-14 0	1	141		4 lb	113
Clams	1	•	1	87	9	Ī	113
Ciams	- 1		1		1		113
Codfish	- 1	- 84	1	79		4 lb filet	113
Crabs	- 1	-400	1	92		is e	113
Duck	- 1	-240	ı	180		14 lb	35
Egg white	- 1	- 52	1	16		white	17
Egg yolk	- 1	-27 0	1	60	- 13	1 yolk	1 -
	- 1		1		1		52
Eggs	- 1	-11 0		82		l, shell removed	113
Frankfurters	- 1	- 93	1	227		2 hoks 1 T	9
Gelatin, dried	- 1		. 1	31		1 T	113
Goose	- 1	- 78		173 81		14 lb filet	113
Haddock	- 1	-12 0	'l	01	- 1	74 in met	
1falıbut		- 9 3	, 1	137	. !	1/2 lb filet	113
Ham (fat)		- 9 5		513		1/4 lb	113
Herring		- 90		154		1 filet	113
Lamb chops		- 9		26		2 chops	113
Lamb, leg		- 9		26	0	1/4 lb	113
2		1					113
Lobster		-38	0		5	⅔ c canned	113
Mackerel		-11	0	20		1 filet	14
Margarine, fortified		0		10		1 T	113
Mince meat		+12		31		14 lb	113
Mutton, leg		- 9	6	21	6	1/4 lb	1
		-		١.	12	6	113
Oysters		-23			52 32	2 chops	113
Pork chops Pork sausage		- 7			56	2 knks	35
Pork sausage Rabbit		-15			98	1/2 lb	113
Salmon		-11			46	1 c canned	113

14

14

MEATS, FISH, AND POULTRY PRODUCTS-(Continued)

	421 - 2		1serage Portion	
Name	Alkaline (+) or lcid (-) Effect	Total Cal- ories	Neasure	Weigh
				grams
Sardines in oil	-11 0	104	4 sardines	50
Scallops	-36 0	84	24 c	113
Shrimp	- 16	72	8	65
Tripe	- 8 1	106	14 lb	113
Tuna m oil	_	111	1/2 c , canned	57
Turkey	-11 0	176	1/4 lb	113
Veal chops	-14 0	236	2 chops	113
Veal cutlet	- 98	203	1/4 lb	113
Whitefish	-10 0	170	34 lb filet	113
	DAIRY	Produc	TS	7
			1	grams
Butter	0.0	103	2 pats	14
Buttermilk	+ 2 2	74	1 glass	210
Cheese, American	- 54	110	1 slice	28 28
Cheese, Cheddar	- 50	110	1 cube	28
Cheese, Cottage	- 45	28	}{ c	_
Cheese, Cream	- 3 4	104	1 cube	28
Cheese, Roquefort	+	109	1 slice	28
Cheese, Swiss	- 50	113	1 slice	28
Cream, sour	l –	29	1 T	15 15
Cream, sweet	00	31	i T	
lce cream, vanilla	+02	214	3∕2 c	100
Milk, chocolate	+15	174	1 glass	210
Milk, condensed (sweetened)		98	2 T	16
Milk, dried skim	+18 0	o7	2 T	16
Vilk, dried whole	+12 0	80	2 T	1
Milk, evaporated	+51	139	14 glass	115
Vilk, goat	i -	147	I glass	210
Vill, human	+	68		100 210
Vilk, skim	+ 2 4	76	I glass	210
Milk, whole (pasteurized)	+ 2 3	145	1 glass	
Whey, dried	+	28	1 T	8
		NUTS	7	
				grams
Almonds	+12 0	88	12 nuts	14
Brazil nuts	+ 4 5	96	2 nuts	14
Buttan	1 ' ' -	95	4 nuts	14

85

26

10 nuts

3 nuts, skin removed

Butternuts

Chestnuts, fresh

Cashew

NUTS-(Continued)

	Alkaline		Irerage Portion	
Name	(+) or lead (-) Effect	Total Cal ortes	Measure	13 e19
Coconut, dried Hazelnuts (filberts) Hickory Peanuts Peanut butter Pecans	+83 +18 -30 -39 -56	79 92 99 83 104 103	2 T 10 nuts 12 nuts 16 nuts, skin removed 1 T 12 meats	gran 14 15 1- 1- 1- 1- 1- 1- 1- 1- 1- 1- 1- 1- 1-
Pistachios Wainuts, black Walnuts English	- 78	93 97	1/2 meats 12 meats	∫ i

	ALECEDIA .	LUCS I U	003	
Apple Pie Chocolate, sweetened Chocolate unsweetened Cocos Codiwer oul Coffee* Corn oul Corn syrup Cottonseed oul Honey Marghe syrup Marmslade orange Wolasses Ohie oul Pickles Potato chips Succotash	+30 +24 +67 +110 +630 00 00 00 -11 + 00 +28 +600 -210 +	346 313 32 27 99 65 99 65 99 80 22 126 56 85 40 99 21 97 94	1 piece, 36 pie 1 T 2 t 1 T 1 T 1 T 1 T 1 T 1 T 1 T 1	grams 100 57 6 6 11 150 11 22 20 14 22 25 14 11 25 17 100
Sucotasas Sugar Granulated (sucrose) Brown Tea* Yeast fresh Yeast dried brewers	0 0 +60 0 +47 0 +17 1	32 19	2 t. 2 t 1 c infusion 1 cake 1 T	8 5 180 14 10

The infusion has little food value except for added sugar and cream. The analytical gures apply to dried collections and details. The infusion has ince took value except for added sugar and cream and figures apply to dried collect bean and dried tea leaves.

These foods are alkaline but because of substances in them which give rise to hippure and in the body they increase the acidity of the urines (see p. 1997).

Ripe olives are soaked in alkali during their manufacture and the alkaline confent. If

apt to be high.

1365 APPENDIX V

V. MAINTENANCE OF ANIMALS FOR NUTRITION EXPERIMENTS

Introduction. Laboratory animals should be housed in well-ventalated rooms free from drufts in order to avoid colds or the blowing around of finely divided dietary dust Rooms should be diffusely lighted during the day and darkened at night, the alternating periods of light and darkness being regular from day to day Exposure of rats (or chicks) to direct sunlight, even though filtered through window glass, must be avoided in vitamin D assay work Rats are nocturnal animals and excessive noise or activity during the day interferes with their normal habits

hir conditioning of rooms is desirable but not essential except where extreme temperatures would otherwise occur The optimum environmen tal temperature for small animals is $24\pm2^{\circ}$ C Provision should be made for screening windows and for frequent washing and flushing of walls and floors, which should be sealed against vermin and wild rodents

Extermination of insects such as roaches, flies, or bedbugs should be commenced as soon as they are observed With sufficient attention and use of modern insecticides, infestation can be effectively eradicated

FLIES AND ROACHES If the room can be cleared of animals, sprayso thoroughly with a potent commercial insecticide such as those containing Pyrethrum, 1 lethane, DDT, methovychlor, or piperouylbutovide Seal room for several hours or overnight Air thoroughly before returning

animals If animals cannot be removed from the room, apply the insecticide by means of a soft brush to walls and floors (especially in corners, eracks, and seams), cage racks, and other equipment except the cages themselves

Bedbugs Paint walls, crevices and equipment with a 5 per cent solution of DDT in kerosene One application may suffice to eradicate a

severe infestation, if not, repeat after six weeks

Excreta should be removed at least every other day Paper rolls or thick layers of newspaper under racks of raised-hottom cages facilitate this operation Hot tin-dipped galvanized wire cages are preferred These sbould be cleaned once every week or two depending upon the number of occupants per cage Cages should be scraped, scrubbed in hot soap solution or soaked in bot trisodium phosphate solution and finally rinsed thoroughly in hot, then cold water

Rats 62 Selectively bred albino or pichatd rats are the animals par

See also Gree man and Duhring The Breeding and Care of the Hbino Rat Philadelphia See also Gree man and Duhring The Breeding and Care of the Hbino Rat Philadelphia Wistar Institute 1931 Extersive box etric data may be found in Donald on The Rat Wistar Institute 1944 Hunt A Laboratory Varual of the Inatomy

⁶⁹ Hand spraying is not nearly as efficient as the use of an electric power spray " Vlany insecticidal sprays contain an insufficient concentration of active pyrethrins A

tery effective product is Fumol made by The Fumol Corporation Long Island City N 1 G For handbooks on the care of laboratory animals and their use in experimental work see Haberland Die operative Tech in der Die Hersperinger 126 Philosope Haberland Die operative Tech in der Die Hersperinger 126 Philosope Assay 2d de Thaladiphas The Blakston Company 1978 Vunch Boostoope Assay 2d de Thaladiphas The Blakston Company 1978 Vunch Boostoope Assay 2d de Thaladiphas The Blakston Company 1978 Vunch Boostoope Assay 2d de Thaladiphas The Blakston Company 1978 Vunch Boostoope Assay 2d de Thaladiphas The Blakston Company 1978 Vunch Boostoope Assay 2d de Thaladiphas The Blakston Company 1978 Vunch Boostoope Assay 2d de Thaladiphas The Blakston Company 1978 Vunch Boostoope Assay 2d de Thaladiphas The Blakston Company 1978 Vunch Boostoope Assay 2d de Thaladiphas The Blakston Company 1978 Vunch Boostoope Assay 2d de Thaladiphas The Blakston Company 1978 Vunch Boostoope Assay 2d de Thaladiphas The Blakston Company 1978 Vunch Boostoope Assay 2d de Thaladiphas The Blakston Company 1978 Vunch Boostoope Assay 2d de Thaladiphas The Blakston Company 1978 Vunch Boostoope Assay 2d de Thaladiphas The Blakston Company 1978 Vunch Boostoope Assay 2d de Thaladiphas The Blakston Company 1978 Vunch Boostoope Assay 2d de Thaladiphas The Blakston Company 1978 Vunch Boostoope Assay 2d de Thaladiphas The Blakston Company 1978 Vunch Boostoope Assay 2d de Thaladiphas The Blakston Company 1978 Vunch Boostoope Assay 2d de Thaladiphas The Blakston Company 1978 Vunch Boostoope Assay 2d de Thaladiphas Delivery of the Williams & Williams & Williams Co. 1931. Men. and B Buolevald Assays Q Philadelphia.

Baltimore The Williams & Williams Co. 1931. Men. and B Buolevald Assays Q Philadelphia.

Baltimore The Williams & Williams Co. 1931. Men. and Baltimore The Williams Co. 1939. Share and Co. 1939. Share and Griffith The Rat in Laboratory Investigation 2nd ed. Philadelphia.

Co. 1939. Farris and Griffith The Rat in Laboratory Investigation 2nd ed. Philadelphia. B Lappincott Co 1949 Farris The Care and Breeding of Laboratory 41 imals New York

excellence for nutrition experiments. Black rats are of course more suitable for study of achromotricbia although white rats develop a condition described as "rusting," comparable to graying of black fur. In their dietary habits rats, like man, are ominiorous. Their period of growth (see Fig. 201) extends over approximately 300 days and their life span is about three years. Thus they live through their cycle at about 30 times the rate of the human species, however, compared with other species.

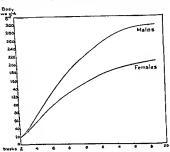


Fig. 294 Normal Growth Curves of Male and Female Albino Rats 14 Food Research Laboratores Breed as Colony

the rate of prepubertal growth of man is slower Rats multiply rapidly and the large litters which they produce permit of a high degree of experimental control. They require but little food and are bence economical to use. For studies of certain nutritional deficiencies to which the rat is immune, notably scurvy, other species of animals must be employed. Variation in species requirements (rat vs. chick) was partly responsible for the discovery of the multiple nature of vitamin D and of the vitamin B complex.

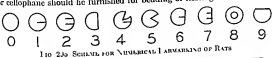
Rats attain sexual maturity at about 70 days but should not be mated for breeding purposes until about 100 days of age. One male may be

of the Rol New York The Wacmillan Co. 1925. See also Howell Anatomy of the Wood Rol Baltimore The Willians & Willians Co. 1326° and especially Greene Anatomy of the Rol I Inhadelphia Amer can I Inhosoph ical Soc ety 1935. "For un form results in vitamina assay work an male stould be bred and reared on

stock dets of constant composition preferably under the direct control of the assay or "These curves are based on 1946 data. Currently (1954) average weights are 20-30 g-ingher from about 10 weeks

A these transformation of the normal growth curve can be sfreeted by platting the lost weight as responsed to the sea seasoning to the method of Zewiter da (Zewiter Hall Young and Zewiter J. Vatration 32 123 (1941) Zewiter da (Zewiter Hall Young and Zewiter J. Vatration 32 123 (1941) Zewiter da (Zewiter Hall Young and Zewiter J. Carry and Addis Am. J. Physical 153 33 (1948)) Thus procedure facilitate comparison of growth data suscent permate supress on of the responses in terms of the slope of the line and it extrepolated asymptotic theoretical maximum) we give

caged with as many as three females at one time. Pregnant females should be isolated before litters are east, unless only one female occupies the cage with the male. The gestation period is about 22 days. The young are usually weaned when three to four weeks old. If the number of pups in a litter exceeds 8, it should be reduced to that number to protect the mother and at the same time produce sturdier rats. If litters are east in wire-mesh cages, the mesh should be less than \(\frac{1}{2}\)2 inch and shredded paper or cellophane should be furnished for bedding or nesting



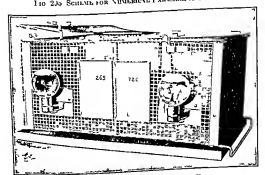


FIG 296 INDIVIOUAL BREEDING CAGE FOR RATS
With Hendrys water fountains removable partition double hd positive
action catch and sliding tray. The front bottom and back are constructed
of a continuous sheet of wire mesh.

 \sim a continuous sheet of wire mesh. Constructed to specifications of Food Re earch Laboratories. Inc. by Norwich Wire Works Norwich N Υ

Rats may be marked by staming the fur with dyes (e.g., methylene blue or pieric acid) by means of an ear tattoo-punch so or by earmarking by means of combinations of mile types of cuts (slits V shaped notches by means of combinations of mile types of cuts (slits V shaped notches or holes) in both ears thus numbering them permanently over a range from 0 to 99 (Fig. 290)

For breeding and stock purposes in small colonies the cage illustrated in Fig 296 may be employed for larger laboratories ricks of suspended drawer type cages are most suitable. These may be equipped to carry outside water bottles and to provide for paper rolls for the collection and removal of excreta (see Fig 297), although stainless-steel trays and

[&]quot; Keeler Science 12, 200 (1940)

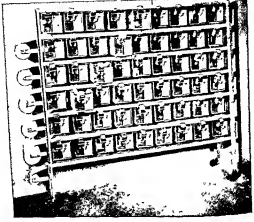


FIG 297 BATTLEY OF RAT CAGES OF THE SUSPENSION TYPE. Manufactured by Norwich Wire Works Norwich, N Y

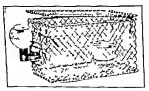


FIG 298 INDIVIDUAL RAT CAGE, OF ALL-WIRE CONSTRUCTION

With Hendryx water fountain positive action catch \onscatter feed cup made of 4-oz out-

ment Jar with plastic cover perforated with 1½ hole Care constructed to specifications of Food Research Laboratories Inc. by Norwich W re Works Yorwich W Y Feed cup supplied by Armattong Cork Co Laboratories and Control of the Property of enster Pa.

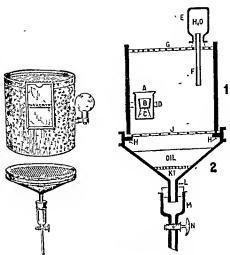


FIG 200 (Left) METABOLISM CAGE FOR SMALL ANIMALS FIG 300 (Right) Cross Section of a METABOLISM CAGE Courtesy Levine and Smith J Leb Clin Ved 11, 168 (1926)

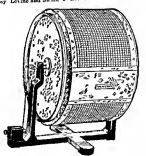


Fig. 301 VOLUNTARY ACTIVITY CASE FOR RATS As used at Yair University

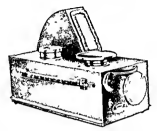


Fig. 302 Exact Weight Shadograph Scale for Weighing Animals Courtery Underwood and Underwood Inc.



FIG 303 TOLEDO SCALE FOR WEIGHING SMALL ANIMALS.
Courtesy Toledo Scale Co Toledo 12 Obio

FOOD RESEARCH LABORATORIES, INC. NEW YORK CITY FOOD CONSUMPTION VITABLE D. ASSAY NO.

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FIG 304, FOOD CONSUMPTION CHART.

FOOD RESEARCH LABORATORIES INC

VITAMIN D ASSAY

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Fig 305 Example of Data Sheet for Vitamin Assay

thick layers of newspaper are in some respects preferable. When they are ready for experiments the rats should be transferred to individual cages of the type illustrated in Fig. 298. These cages are of all wire construction of the type illustrated in Fig. 298. These cages are of all wire construction of the have false bottoms to minimize access to excreta 65. They are easily attributed.

Specially prepared diets should be fed in a nonscatter type of food cup to permit the collection of more accurate data on food consumption Suitable for this purpose is a 3- or 4-oz screw cap ountment jar with a

[&]quot;Cages and accessories may be obtained from the Norwich Wire Works Norwich New York the Goe H. Wahmann Mig Co Baltimore Md or the A B Hendryx Co New Haven Connections.

Rats may be weighed on a spring bilance (Chatillon or Hanson), but for more exact work an Exact Weight or Toledo scale of 250-500-, or 1000-g capacity, sensitive to 1 g should be used (See Figs 302 and 303) These bilances are also suitable for weighing food cups Records are kept of the weekly or semiweekly weighings on form sheets such as are illustrated in Figs 304 and 305 A modified type of growth chart (Fig 306) in which the time coordinates (absense) are divided into weeks and days has proved very useful

Breeding and Stock Diets. When rats are reared for purposes not requiring careful control of their nutrition, they may be fed a mixed diet requiring careful control of their nutrition, they may be fed a mixed diet consisting of clean table scraps, bread and milk, or dog biscuit⁵⁷ supplemented with milk or fresh vegetables such as carrots or lettuce, once or twice a week Specially prepared rations for rats and other laboratory animals are available on the market and have proved quite satisfactory ⁵⁵ Mixed poultry rations (growing mash) or calf meal may also be used For Mixed poultry rations (growing mash) or calf meal may also be used For Mixed poultry rations (growing mash) or calf meal may also be used For Mixed poultry rations (growing mash) or calf meal may also be used for the special poultry rations of rats are defined by the special poultry rations of the fat-soluble vitamins be fed diets which limit or control their reserves of the vitamins in question

I FOOD RESEARCH LABORATORILS' RAT STOCK DIET*9

I FOOD ICESEAROR			
Whole wheat hellow corn Nonfat milk solids Hydrogenated cottonseed oil Veat meal (65% protein) Alfalfa (hiln-dried) heast (dried)	10 20 15	Vitamin B-complex mixture ⁵⁰ Vitamins in oil ⁵¹ Cellulose powder NaCl MnSO ₄ H ₂ O Pork liver (vacuum dried)	1 1 0 5 0 02 0 48

⁴ The dog biscurts used should be of known nutritional quality as there is a great variation among them some being incapable even of supporting normal growth Certain brands ton among them some being incapable even intuitives or mashes.

New City New York.

10 Based originally on the Sherman Diet B (J Biol Chem 53 49 (1922) 60 5 (1924))

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18 Based original Properties of Sherman Diet B (1924)

18 Based original Properties of Sherman Diet B (1924)

Thamine Recomplex mixture Thamine Rhofavine Pyridoxine Niacin Calcium pantothenate p-Aminobenzoic acid Inostol	1 g contain 0 6 mg 1 2 mg 0 4 mg 5 0 mg 4 0 mg 2 5 mg 100 mg	ss Cholme chloride Liver concentrate 1 20 Biotin Folio acid Cyanocobalamin Cellulose powder q s	200 mg 25 mg 1 µg 1 µg 1 µg 1 g
--	---	---	--

71 Vitamins in oil 1 g contains

g contains	200 units
Vitamin A	20 units
Vitamin D	12 mg
α Tocopi erol	100 µg
Menadione	1 g
Cottonseed oil q.s.	

of pet foods are available in the form of unpelleted mixtures or mashes as the form of compressed a Ralston Purina Co. St. Louis Mo. supplies such rations in the form of compressed the Ralston Purina Co. St. Louis Mo. supplies such rations in the form of compressed the Ralston Purina Co. St. Louis Mo. supplies such rations are available through Rockland Farms guines, pigs, rabbits dogs etc.) Similar rations are available through Rockland Farms guines, pigs.

2 BILLS MODIFICATION OF STEENBOCK STOCK DIET	THE	Pork liver (vacuum dried) Calcium carbonate	0.5
Yellow corn Dried whole milk	57 25	Sodium chloride Lettuce or spinach about 5	
Linseed oil meal	12 3 7	4 F R. L VITAMIN-RESTRICTED	DIET
Crude casein		Wheat meal (entire kernel)	37 0
Alfalfa leaf meal	15	Wheat mear (critic kerner)	37 0
Iodized table salt	0 4	Yellow corn meal (entire kernel)	22 0
Calcium carbonate	0 4	Whole milk powder74	10
3 A. D M A BREEDER D	157	Dred yeast**	0 2
3 A. D. A. A. DREEDER D.		Dried pork liver	20
Wheat meal (entire kernel)	33 Q	Dried alfalfa	0 0
Yellow corn meal (entire kernel)	34 0	Calcium earbonate	0.5
Whole milk powder	21 0	Sodium ehloride	0.02
Old process linseed oil meal	7 0	Manganese sulfate	
Alfalfa leaf flour (green)	20	Liver concentrate (Wilson 1 20)	0 1

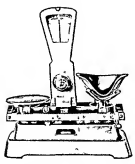


FIG 307 EXACT WEIGHT SCALE FOR PREP-ARATION OF EXPERIMENTAL DIETS Courtesy Underwood and Underwood, Inc

Typical diets which have proved successful are shown on p 1373.4 These may be prepared by means of scales of the type illustrated in Fig. 307 Salt Mixtures Synthetic diet mixtures used in nutritional research on rats include special combinations of inorganic salts whose composition

¹³ Bills Honeywell Wirick and Nussmeier J Biol Chem 90 619 (1931) Landow Peterson and Steer bock J Biol Chem 84, 419 (1929)

¹¹ In some laboratories a more highly restricted duet may be used with safety in vitamia A work. For this purpose skim milk powder may be substituted for whole milk powder is Not irraduated or fortified with vitamin D

is based on the analysis of their milk 5 or urine The original Osborne-Mendel and McCollum-Davis salt mixtures, given below, have been modified to include readily available commercial forms of the chemicals employed (see p. 1262). The modification of the Osborne-Mendel mixture by Hawk and Oser includes salts which yield a similar stoichiometric mixture but, by avoiding acids and carbonates, eliminates the need for evaporation and deligidration.

OSBORNE MENDEL SALT MIXTURE 6

h ₂ CO ₂ 1	134 8 H ₄ PO ₄ 141 3 HCl 24 2 H-SO ₄ 34 2 Citric acid + H	103 2 Fe citrate 1 53 4 NaF 9 2 KAl(SO ₄) ₂ 10 111 1 MnSO ₄ KI	0 062 0 0245 0 079 0 020
----------------------------------	---	--	-----------------------------------

All amounts are given in grams. The acids are added to the carbonates and ferne citrate, and stock solutions of the other salts are then added in proper amount. After effer escence is completed the mixture is dried at 90° to 100° C and ground to a fine powder. These amounts of phosphone, hydrochloric and sulfure acids are equivalent to 71, 121, and 5 2 ml, respectively, of the concentrated acids. See also p. 1262.

HUBBELL, MENDEL, AND WAREMAN SALT MIXTURE 78 HAWL OSER SALT MIXTURE NO 311 543 0 CaCOa 308 2 Ca Citrate 4H.O 25.0 MgCO: 112 8 Ca(H,PO4), H2O 16 0 MgSO. 218 7 L.HPO. 69 O NaCl 124 7 KCI 112 0 77 0 KCI \aCl 212 0 KH.PO. 68 5 CaCO. FePO, 4H2O 20 a 35 1 3 MgCO, Mg(OH), 3H,O 0.08 38 3 KT MgSO, anhydrous 0.35 MoSO. 91 41 Fe NH, Citrate USP 1 00 NaF _b 98 CuSO, 5H2O 0.17 AL(SO,),K SO. 0 76 NaF 0 90 CuSO. 16 7 1 07 MnSO, 2H2O 0 54 KAI(SO4), 12H2O 0 24 Κī 1000 00 100 00

Mr Collum Davis Salt Mixture No. 18579

STEENBOCK SALTS 4060

Ca lactate Ca(II;PO ₄) ₂ II;O I ₂ :HPO ₄ NaH;PO ₄ II;O ACI MgSO ₄ (anhy drous) Fe citrate	35 15 \aCl 14 60 \UgSO ₄ 7H ₂ O 25 78 \aghtarrow 9 38 \aghtarrow 467 \cantrow 7 19 \cantrow 3 19 \text{FeCaH}_2O_2\text{1}_2O 3 19 \text{FeCaH}_2O_2\text{1}_2O 4 67 \text{Alpha}_2\text{1}_2O 4 67 \text{Alpha}_2\text{1}_2O 4 19 \text{FeCaH}_2O_2\text{2}_2O 4 19 \text{Alpha}_2\text{2}_2\text{1}_2O 4 19 \text{Alpha}_2\text{2}_2\text{1}_2O 4 19 \text{Alpha}_2\text{2}_2\text{1}_2O	233 6 246 0 358 0 696 0 698 0 194 0 99 8 1 6
--	--	---

Guinea Pigs (Cavies). The most suitable guinea pig for nutrition work is the English short smooth-haired variety, whose manifold coloring facilitates identification. Not all pure white guinea pigs are albinos since they may have dark-colored instead of pink eyes.

Guinea pigs must he housed away from drafts in dry eages, preferably with false hottoms although beds of sawdust or shavings are satisfactory. Dry hay should be provided at all times. They are herhavorous and may be fed clean hay and grain (oat meal, bran, wheat middlings, etc.) supplemented with fresh vegetables (carrots, beets, alfalfa, lettuce, celery, etc.)

Most vegetables must be sound and wholesome because guinea pigs will reject spoiled food. Diets containing fats (e.g., codiiver oil) must not be fed when rancid. The addition of dired yeast to the grain mixture is frequently desirable. Specially prepared diet mixtures for guinea pigs are available commercially. Water should be fed spannigly when fresh vegetables are supplied.

Most investigators purchase guinea pigs on the market, since their nutritional listory is seldom of importance. However, they should be purchased from a reputable breeders since they are prone to earry pul monary and other infections when raised or shipped under adverse conditions. Guinea pigs breed about once every three or four months, the gestation period being from 65 to 70 days. Mating should commence not before the age of three months. The lactation period should last about two to three weeks.

White Mice. The white mouse has been employed with some success in studies in nutrition. Advantages of the mouse include a smaller food requirement, more rapid growth, and frequency of reproduction. The dietary needs of the mouse are similar to those of the rat although a somewhat higher protein content is recommended.

Advantages of the rat over the mouse for nutritional work include the fact that the rat has been more extensively studied, thus affording more norms for comparison, also at is hardier than the mouse, and thus not so prone to contract minor illnesses or to be adversely influenced by changes in temperature, etc

It has been claimed that mice require vitamin C for normal development 84

Dogs. 55 The feeding of a purely "synthetic" food mixture for long periods, in metabolism tests using dogs as experimental animals, is a procedure for the successful accomplishment of which investigators in metabohsm have long striven It remained for Cowgill to solve this important problem in dietary technique. His work was an extension of that initiated by Karr 86

The principles upon which the procedure is based are that the diet consist of a mixture of isolated food substances which supply the body with all factors necessary for proper untrition, except a single dietary factor, and that the variable nutritive factor shall be administered apart from the diet mixture. In this discussion vitamin B (complex) is regarded as this variable nutrition factor

In a test where vitamin B is the variable factor, the diets7 should contain (1) protein, proper in kind and amount, \$4 (2) carbohydrate, \$9 and fat 90 to furnish satisfactory energy, (3) mineral salts, 91 proper in kind and

[&]quot;See footnote 92 p 1378 11 Cowgill suggests the use of one of the following salt mixtures

SALT VIXTURE (MARE		SALT MIXTURE (COWGILL)	
NaCl Ca lactate Mg citrate Fe citrate Iodine in KI (Lugol s solution)	10 g	NaCl Vig citrate KiisPO: CaHPO: 2H:O KCI Fe citrate KI	3S 0 32 5 12 2 7 8 7 0 1 8 0 5

Karr s salt mixture when fed with bone ash as a source of phosphate on the basis of 0.2 g and 0.4 g. respectively per hio unit of food has given successful results during periods lasting over five months

Where 0.4 g of agar agar per kilo unit is used as a source of roughage Cowgill scalt the deep of agar agar per sale of 03 g per kilo unit of food it is supposed to furnish the deep of the the dog the various essential morganic substances equal to or slightly in excess of that one the various essential morganic amendates close to a significant descriptions of the second and the second a

Millemer and Thuber Food Research 1 399 (1936)

^{**} Cowgill J Biol Chem 56 725 (19°3) This procedure has been used at Yale Univer sity for periods of five months and over with excellent results

[&]quot;Karr J Biol Chem. 44 200 (1920)
"Calculation of the Diet. Construct the diet on a lg of body weight hasis or form what might be called a kilo unit of food Done weathing 7 kg and over require for the maintenance might be called a kilo unit of food Done weathing 7 kg and over require for the maintenance of body weight between 70 and 80 Calories per kg per day 80 Calories permits maint of body weight between 70 and 80 Calories per kg per day 80 Calories per kg. tenance in adult animals and allows a slight increase of body weight in growing dogs Dogs weighing considerably less than 7 kg require slightly more energy intake in order to main tan their body weight this fact however does not invalidate the choice of 80 Calories as the kilo unit quantity Any variation in energy intake from this figure due to the size of the animal will not seriously affect the intake of nitrogen and salt mixture if liberal quantities such as 0 S g N and at least 0 2 g salt mixture are furnished with every kilo unit of food

For further discussion of the hasis for this diet consult the original paper This may be casein extracted meat (from the Valentine Meat Juice Co Richmond

Va) or commercial coag lated egg albumin Casein is to be preferred

^{...} commercial coog lated egg albumna tasenh is to be picasired.

Sucrose is a desirable source of carbohy drate pure starch or destrin make the diet

of pasty when mostemed by salva W ole cereals however are suitable

amount, (4) sufficient vitamin A, 22 (5) roughage, 22 and (6) water 24 The energy requirement as shown in the table of the "kilo unit" below 15 somewhat high for dogs whose activity is restricted, e.g., in metabolism cages. In such cases it is advisable to reduce the calories to 60-70 per kg body weight by substituting sucrose for land. The vitamin B (the variable

hilo Unit of Casein Food*

	Amount	Calories	Percentage
Casem (81 9 per cent pure, 12 7 per cent \) Sucrose Lard Butterfat Bone ash Salt mixture† Totals	g 6 3 5 84 2 83 1 17 0 40 0 20 16 74	20 8 23 4 25 5 10 5	37 6 34 9 17 0 7 0 2 3 1 2

^{*} This kilo unit contains 80 calories 45 per cent of which are furnished by fat and 08s of hitrogen

† See footnote 91 p 1377

factor) should be fed separately from the rest of the diet, in the form of dred stast (about 0.4 g per kilo unit of food) or preferably a synthetic vitamin mixture, or the various components of the vitamin B complex may be fed as separate supplements Excessive amounts of sugar in the diet of dogs will cause diarrhea, hence cereals (but not pure starch) are better sources of carbohydrate. It has been claimed that meat is an indispensible article of diet in tests on dogs. As a matter of fact, meat in the feeding of dogs is not essential if the diet is adequate with respect to protein and other nutrents.

Cowgill's work shows that "monolony of diel is not per se the cause of failure of appetite loss of the desire to eat is rather an expression either (1) of the failure of the food being fed to nourish the animal properly, or (2) of an adjustment of the dog to its energy requirement when offered more than is necessary of a food mixture that is adequate in all respects"

 U_2

VI. ANALYSIS OF VARIANCE "

Table 1

ANALYSIS OF VARIANCE (ASSAY NO 34005)

S.

Sample Vitamin A Concentrate Number of groups 4 (2 assay and 2 reference)

 S_1

Assay level 217 200 USP u/g Number of rats per group (N) 9 males

 U_1

Dose (mg)	0 88 (1 5 u)	1 47 (2 5 u)	0 00691 (*15 u)	0 01154 (25 u')
Totals Weans	28 Y ₁ 31 Y ₂ 36 35 30 10 34 30 24 208 (e) 28 7	20 49 65 37 30 37 36 35 41 350 (/)	24 32 27 31 22 34 26 29 13 238 (g) 26 4	45 68 38 27 50 45 58 32 26 Y ₂ 389 (h) 43 2
	of sums of squ +f+g n lows \times Columns Columns (betw	reen doses) = -	$0 + 238 + 389)^{2} = 0$ 9×4 $1 + P + q^{2}$ $1 + Rows$ $58^{2} + 350^{2} + 238^{2} + 97029$ $1 + C$ $36^{2} - C$ $26^{2} - C$	$\frac{389^2}{2} - C$

 $= 28^{2} + 31^{2} + 36^{2}$ =47,791-42,36736Experimental error = total - columns = 5423 64 - 1746 97 = 3676 67

[&]quot;This is included to illustrate a useful experience at lesign for boassays. For references to important text-books on biometries see footnote 42 p 1130

amount, (4) sufficient vitamin A, 22 (5) roughage, 22 and (6) water 34 The energy requirement as shown in the table of the "kilo unit" below is somewhat high for dogs whose activity is restricted, e.g., in metabolism cages In such cases it is advisable to reduce the calones to 60-70 per kg body weight by substituting sucrose for lard. The vitamin B (the variable

KILO UNIT OF CASEIN FOOD*

	Amount	Calories	Percentage
Casein (819 per cent pure, 127 per cent \) Sucrose Lard Butterfat Bone ash Salt mixture? Totals	g 6 3 5 84 2 83 1 17 0 40 0 20	20 8 23 4 25 5 10 5	37 6 34 9 17 0 7 0 2 3 1 2

^{*} This kilo unit contains 80 calories 40 per cent of which are furnished by fat and 0 85 of nitrogen

† See footnote 91 p 1377

factor) should be fed separately from the rest of the diet, in the form of dried yeast (about 0.4 g per kilo unit of food) or preferably a synthetic vitamin mixture, or the various components of the vitamin B complex may be fed as separate supplements Excessive amounts of sugar in the diet of dogs will cause diarrhea, hence cereals (but not pure starch) are better sources of carbohy drate. It has been claimed that meat is an indepensable article of diet in tests on dogs. As a matter of fact, meat in the feeding of dogs is not essential if the diet is adequate with respect to protein and other nutrents.

Cowgil's work shows that "monolony of diet is not per se the cause of failure of appetite loss of the desire to cat is rather an expression other (1) of the failure of the food being fed to nourish the animal properly, or (2) of an adjustment of the dog to its energy requirement when offered more than is necessary of a food mixture that is adequate in all respects"

ah

VI. ANALYSIS OF VARIANCE 95

Table 1

ANALYSIS OF VARIANCE (ASSAY NO 34005)

Assay level 217,200 USP u/g Sample Vitamin A Concentrate Number of rats per group (N) 9 males Number of groups 4 (2 assay and 2 reference)

	S_1	S,	<i>U</i> ₁	<i>U</i> :
Dose (mg)	0 88 (1 5 u)	1 47 (25 u)	0 00691 ('1 5 u ")	0 01154 ('25 u")
Totals	28 Y ₁ 31 Y ₂ 36 35 30 10 34 30 24 258 (e) 28 7	20 49 65 37 30 37 36 35 41 350 (f) 38 9	24 32 27 31 22 34 26 29 13 238 (g) 26 4	45 08 38 27 50 45 58 32 26 Y _n 389 (h) 43 2

Calculation of sums of squares
$$C = \frac{(e+f+g-n)^2}{\text{Rows} \times \text{Columns}} = \frac{(258+350+238+389)^2}{9\times 4} = \frac{(1235)^2}{36} = 42367\ 36$$

$$\text{Columns (between doses)} = \frac{e^1+f^2+g^2}{8\text{Rows}} - C$$

$$= \frac{258^2+350^2+238^2+389^2}{9} - C$$

$$= \frac{397,029}{9} - 42,367\ 36 = 1746\ 97$$

Total =
$$Y_1^2 + Y_2^2 = Y_1^2 - C$$

= $28^2 + 31^2 + 36^2 = 26^2 - C$
= $47,791 - 42,367 \cdot 36$
= $5423 \cdot 64 = 1746 \cdot 97 = 3676 \cdot 1$

Experimental error = total - columns = 5423 64 - 1746 97 = 3676 67

This is included to illustrate a useful experimental design for bioassays. For references to important text books on biometrics see footnote 42 p 1130

Talle 2 FACTORIAL ANALYSIS OF VARIANCE BETWEEN DOSES

Variations	Fact	orial Co	ficients	(x)	Duisor	Sum of Prod-	Variance (ExYp)*
Due to	s,	S ₂	U ₁	U ₁	\ 2x1	ucts ExYp	$\frac{(2x^2p)}{\lambda 2x^2}$
Difference be-	-1	-1	+1	+1	36*	+19†	10 03
Slope of dose re-	-1	+1	-1	+1	36	+243	$1640 \ 25 = B^2$ $40 \ 50 = B$
Departure from parallelism	+1	-1	-1	+1	36	+59	96 69
Total (Yp)	258 (e)	350 (f)	238 (g)	389 (h)			

Note D and B take the again of the bunns of Products from which they are derived $^{\circ}9$ ((-1)² + (-1)² + (-1)³ + (+1)³ + (+1)³) = 36 (-1)²

Table 3 ANALYSIS OF VARIANCE

Variations Due to	Degrees of Freedom (DF)	Sum of Squares (SS)		Variance	Signifi- cant F*
Difference between samples Slope of dose-response cur Departure Irom parallelism Experimental error Total	e 1	10 03 1640 25 96 69 3676 67 5423 64	10 03 1640 25 96 69 114 90 (st)	0 09 14 28 0 84 1	4 15 4 15 4 15

 $s \approx \sqrt{11490} = 1072$

^{*} From Suedecor Statistical Methods Applied to Experiments in Agriculture and Biology. 4th ed Ames lows Collegiate Press 1946 p 224 See also footnote 42 p 1130

Table 4

CALCULATION OF LITTMATE OF POTENCY

$$M = \log \frac{\text{potency of } U}{\text{potency of } S} = \frac{KID}{B}$$

 $M = \log \frac{\text{potency of } U}{\text{potency of } S} = \frac{KID}{B}$ K = 1 (for two-dose assays, for three-dose assays K = 1 635)

$$I = \log \frac{\text{large dose}}{\text{small dose}} = \log 1 667 = 0 2219$$

D = 3 17 B = 40 50 from factorial analysis (Table 2, on p 1379)

$$M = \frac{1 \times 0.2219 \times 3.17}{40.50} = 0.0174$$

Log estimated potency of sample (LP) = log of assumed potency of sample (AP) + MLog(AP) = log 217,200 = 53369

$$V=0.0174$$

 $\log (EP) = \overline{53543}$ antilog 5 3513 = 226,100 = Estimated potency of sample in vitamin A units per gram

Table 5

CALCULATION OF ERROR OF ESTIMATE

$$S_M = \log \text{ standard error} = \frac{sKI \sqrt{B^2 + D^2}}{B^2}$$

where

$$K \approx 1$$

$$I = 0.2219$$

$$B^2 = 1640.2$$

$$B^2 = 1640 25$$

 $D^2 = 10 03$ from factorial analysis (Table 2, above)

from factorial analysis (1406 2, 4650 28)
$$S_M = \frac{10.72 \times 1 \times 0.2219 \times \sqrt{1650.28}}{1640.25} = 0.0589$$

 $\log (EP) \pm S_M = \log (limits of potency)$

$$5\ 3543\ -0\ 0589\ =\ 5\ 2954$$

Standard error as per cent of estimate = 13 6 per cent

VII. FOUR-PLACE TABLE OF LOGARITHMS

	VII. FOUR-PLACE TABLE OF							<u> </u>		Proportional Parts												
Natural Numbers	0	1	.	2	3		4	5		6	7	8	9	1	2	3	4	5	б	7	8	9
10 11 12 13 14													0374 0755 1106 1430 1732									37 34 31 29 27
15 16 17 18 19	204 230 25 27	1 2 14 2 53 2 88 2	068 330 577 810	200 23: 260 28:	5 21 10 23 11 26 33 28	22 80 25 56	2149 240 264 287	217 524 826 829	30 72 00	245 245 269 292	222 248 271 271 294	0 250 8 274 5 296	7 2014 3 2279 4 252 2 276 7 298	2 2 2	6 5 5 5 4	1 2	10	12	11	111111111111111111111111111111111111111	21 20 5 19 5 11	25 24 22 22 21 5 20
20 21 22 23 24	30 32 34 36 38	103 223 243 17 02	032 3243 344 363 382	30: 32: 34: 36: 36: 36: 38:	54 30 63 33 64 3 55 36 38 3	175 184 183 674 856	309 330 350 369 387	631 133 235 237 438	18 24 22 11 92	313: 334: 354 372 390	9316 336 1350 9374 6392	0 318 5 338 0 357 7 376 7 394	1 320 340 9 359 6 378 5 396	1 2 2 2 2 2 2	4		5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5	3 1	0 1:	2 1 2 1 1 1 1 1	4 1 4 1 3 1 2 1	7 19 6 18 5 17 5 17 4 16
25 26 27 28 29	4 4 4	50 14 72	416 433 448	6 41 0 43 7 45	83 4 46 4 02 4	200 362 518	421 437 453	6 42 8 43 3 45	232 333 48	124 140 156	9 420 9 442 4 457	55 128 5 114 9 159	16 413 31 429 40 445 4 460 12 475	8 2 6 2 9 2	3		5	77656	8 1			4 15 3 15 3 14 2 14 2 13
30 31 32 33 34	4.5.55)14)52 183	192 506 519	8 49 5 50 8 5	142 4 179 5	953 097	496 510	9 4 39 5	983 119 210	499 513 520	7 50 2 51 3 52	11 502 15 518 16 52	30 490 24 503 59 517 89 530 16 542	8 1	3333	ŀ	1	6655	- 1	- 1	01 91 91 91	1 13 1 12 1 12 0 12 0 11
35 36 37 38 39		\$63 682 798 911	55 56 58 59	75 5. 94 5 09 J 22 5	587 705 821 933	559 571 583 534	9 56 7 57 2 58 1 59	115 295 435 555	623 740 853 960	56. 57. 58. 59.	35 56 52 57 56 58 17 59	47 56. 63 57 77 58 88 59	39 553 58 567 75 578 88 58 39 001	0 6 9 0		222	44333	5 5 5 4	6 6	171-1-1-1	ŝ.	0 11 0 11 9 10 9 10 9 10
40 41 42 43 44		333. 343.	63 64	13,6 15,6 11,6	253 355 454	626 636 646	3 62 5 63 4 64	74 6 75 6 74 6	28 38 48	63 63	94 63 95 64 93 65	04 63 05 64 03 65	07 61 1 12 62 1 14 63 1 15 64 1 13 65 1	25 25 22		2	3	4444	5 5 5 5	1	1 1 1 1 1 1	9 10 8 9 8 9 8 9
45 46 47 48	9	030	200	"	0920	103,	ω _l v:	*34 p	034	963	သမေ	10-1 02	09 68 02 67 94 68 84 68 72 69	81	11	2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2		4	5 3 4	5	1	988888
5 5 5 5	0 1 2 3	1707	616	0.841	7033	1710	1157	110	711	<u>s</u> [71	20171	25 71	159 70 143 71 126 72 108 73 188 73	521	. 1	2 2 2 2 2	3 2 2 2	333333	111111	50055	66666	88777

APPENDEX VII

FOUR-PLACE TABLE OF LOGARITHMS-(Continued)

		r	0016-1	LACE	IAL		. ~-							_	_	_			
		!										P_{I}	оре	rtt	one	al I	Par	is	
Vatural Sumbers	0	1	2	3	1	5	6	7	8	9	1	2	3	1	5	6	7	8	9
55 56 57 58 59	7404 7182 7559 7631 7709	7412 7490 7566 7642 7716	7419 7497 7574 7649 7723	7 127 7505 7582 7657 7731	7435 7513 7589 7664 7738	74 13 7520 7597 7672 7745	7451 7528 7604 7679 7752	7459 7536 7612 7686 7760	7466 7513 7619 7694 7767	7474 7551 7627 7701 7774	1 1 1 1	2 2 2 1 1	22222	33333	1111	5 5 5 4 4	55555	6 6 6	777777
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